

Substance Name: 2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec-butyl)phenol (UV-350)

EC Number: 253-037-1

CAS Number: 36437-37-3

MEMBER STATE COMMITTEE

DRAFT SUPPORT DOCUMENT FOR

2-(2H-BENZOTRIAZOL-2-YL)-4-(TERT-BUTYL)-6-(SEC-BUTYL)PHENOL (UV-350)

Presented by the dossier submitter at MSC-30

14 June 2013

(Not concluded by MSC)

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Substance Name: 2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec-butyl)phenol (UV-350)

EC Number: 253-037-1

CAS number: 36437-37-3

The substance is identified as vPvB according to Article 57 (e).

Summary of how the substance meets the criteria set out in Article 57(e) of REACH

Persistency:

According to a weight-of-Evidence argumentation UV-350 has to be considered vP and therefore also P.

Overview of the conclusions of the weight-of-evidence approach:

- QSAR calculations indicate that UV-350 is a borderline case for meeting the screening criterion for persistence. Also, the structural similar substances UV-320, UV-327 and UV-328 all show a low biodegradability in ready biodegradation tests. Therefore in a read-across it is plausible that this will be the case for UV-350 as well.
- Read-across assessment on EC 407-000-3 and its first metabolite: Very slow dissipation in aerobic systems (sediment and water) near or above the vP-trigger value based on data for the different compartments with and without temperature correction. Modelling of anaerobic system shows a DegT50 > 180 days already at 20°C. Degradation of the substances in question has to be even longer;
- For UV-327 and UV-328 there are monitoring studies available showing that the substances were found decades after environmental exposure has stopped. Model calculations indicate that these findings can only be explained if the $DegT_{50}$ is larger 180 days.
- Further supporting information:
 - Simulation of the complex degradation pathways gives a mechanistic explanation for similarities and findings;

Thus, applying the weight-of-evidence approach the substance fulfills the P and the vP-criterion of REACH Annex XIII

Bioaccumulation:

Based on a MITI-BCF study the substance fulfils the B and the vB criterion of REACH Annex XIII.

Conclusion:

In conclusion UV-350 meets the criteria for a vPvB-substance according to Article 57 e).

Registration dossiers available: No

Note: This dossier is one of four dossiers for the SVHC-identification of several phenolic benzotriazoles as vPvB-substances and in two cases also as PBT-substances. Since these substances are structurally very similar and relevant data on individual substances for some endpoints is scarce, in these instances all information for all four substances of the set is given to allow an assessment based on read-across and a weight-of-evidence-approach in an analogue approach. All relevant available experimental data on the substances in question is presented in a read-across-Matrix in Annex 1. In the individual chapters only the relevant data for assessing the individual endpoint will be presented. The set of the four phenolic benzotriazoles composes of:

Name	EC-nr.	CAS-nr.	Trade name used in this dossier	Structure
2-benzotriazol-2-yl- 4,6-di-tert-butylphenol	223-346-6	3846-71-7	UV-320	OH N
2,4-di-tert-butyl-6-(5- chlorobenzotriazol-2- yl)phenol	223-383-8	3864-99-1	UV-327	
2-(2H-benzotriazol-2- yl)-4,6- ditertpentylphenol	247-384-8	25973-55-1	UV-328	OH N N
2-(2H-benzotriazol-2- yl)-4-(tert-butyl)-6- (sec-butyl)phenol	253-037-1	36437-37-3	UV-350	

Table 1: Overview of the phenolic benzotriazoles proposed for SVHC-identification.

JUSTIFICATION

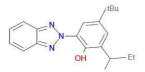
1 Identity of the substance and physical and chemical properties

1.1 Name and other identifiers of the substance

Table 2:	Substance	identity
	Jubstance	lucificity

EC number:	253-037-1
EC name:	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec- butyl)phenol
CAS number (in the EC inventory):	36437-37-3
CAS number:	36437-37-3
Deleted CAS numbers:	122245-62-9, 142513-61-9, 153613-76-4, 188025-37- 8, 189456-67-5
CAS name:	Phenol, 2-(2H-benzotriazol-2-yl)-4-(1,1- dimethylethyl)-6-(1-methylpropyl)-
IUPAC name:	2-(2H-Benzotriazol-2-yl)-6-sec-butyl-4-tert- butylphenol
Index number in Annex VI of the CLP Regulation	-
Molecular formula:	C ₂₀ H ₂₅ N ₃ O
Molecular weight range:	323.432 g/mol
Synonyms:	2-(2-Hydroxy-3-sec-butyl-5-tert-butylphenyl) benzotriazole;
	2-(2-Hydroxy-3-sec-butyl-5-tert-butylphenyl)-2H- benzotriazole;
	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec- butyl)phenol;
	2-(3-sec-Butyl-5-tert-butyl-2- hydroxyphenyl)benzotriazole;
	2-(3'-sec-Butyl-5'-tert-butyl-2'- hydroxyphenyl)benzotriazole;
	4-tert-Butyl-6-sec-butyl-2-(2H-benzotriazol-2- yl)phenol;
	Chisorb 350;
	Eversorb 79

Structural formula:



1.2 Composition of the substance

Name: 2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec-butyl)phenol

Description: mono-constituent

Degree of purity: \geq **98** %¹

Table 3: Constituents

As this substance is a monoconstituent substance this information in not relevant.

Constituents	Typical concentration	Concentration range	Remarks
2-(2H-benzotriazol-2- yl)-4-(tert-butyl)-6- (sec-butyl)phenol	≥ 98 %	≥ 95 - 100 %	
EC-Nr. 253-037-1			

Table 4: Impurities

Impurities	Typical concentration	Concentration range	Remarks
n.a.			

Table 5: Additives

Additives	Typical concentration	Concentration range	Remarks
n.a.			

¹ From C&L notifications

1.3 Physico-chemical properties

Property	Value	Remarks
Physical state at 20°C and 101.3 kPa	-	-
Melting/freezing point	81 – 83 °C	Rosevear, Judi; Australian Journal of Chemistry 1985, V 38(8), P1163-76 CAPLUS
Boiling point	458.0±55.0 °C	calculated properties using Advanced Chemistry Development (ACD/Labs) Software V11.02 (©1994- 2010 ACD/Labs)
Vapour pressure	5.22*10 ⁻⁹ Torr, T= 25 °C	calculated properties using Advanced Chemistry Development (ACD/Labs) Software V11.02 (©1994- 2010 ACD/Labs)
Water solubility	0.1395 mg/l T= 25 °C,	QSAR estimation from log KOW with the EPISuite module WSKOW v1.41; log Kow used for calculation: 6.31 (also estimated, see below)
Partition coefficient n- octanol/water (log value)	6.951 ± 1.251 T = 25 °C 6.31 7.11	calculated properties using Advanced Chemistry Development (ACD/Labs) Software V11.02 (©1994- 2010 ACD/Labs) according to SCIFINDER data EPISuite v.4.10 COSMOtherm v. C30_1201
Dissociation constant	-	-
[enter other property, if relevant, or delete row]	-	-

Table 6: Overview of physicochemical properties

2 Harmonised classification and labelling

No harmonised or agreed classification is available for the substance. Therefore the self classifications according to Regulation (EC) 1272/2008 (CLP) from ECHA's C&L Inventory database (accessed 09.10.2012) are provided in Annex 2 to give some indications on the hazards of the substance.

3 Environmental fate properties

3.1 Degradation

3.1.1 Abiotic degradation

3.1.1.1 Hydrolysis

The chemical bond between the benzotriazole group and the aromatic ring is generally expected to be very strong and also able to withstand degradation due to hydrolysis (see also 3.1.2.1.1) and also the aliphatic groups in the side chains of the phenol ring are functional groups that are expected to be generally resistant to hydrolysis. Due to the high log KOW and the high adsorption potential to organic carbon the substance will adsorb to sewage sludge and suspended organic matter when it is released to the sewage treatment system respectively to the aquatic environment.

Therefore hydrolysis is not expected to be a relevant pathway of elimination of UV-350.

3.1.1.2 Phototransformation/photolysis

Phenolic benzotriazoles are mainly used as an UV-absorber. This means that on the molecular level UV-radiation excites the phenolic benzotriazole. In this excited state a proton from the OH-group is transferred to a nitrogen atom. From this structure a radiationless deactivation coupled with another proton transfer from the nitrogen back to the OH-group will bring the molecule back into its ground state. The UV-protection properties are based on this fully reversible and non-destructive process. Therefore photolysis can be regarded as a negligible degradation path, nevertheless the different compartments will be briefly discussed.

3.1.1.2.1 Phototransformation in air

An estimation for atmospheric degradation with OH-radicals has been conducted with AOPwin v1.91 (US EPA, 2011) assuming a 12 hour-day and a OH-concentration of $1.5*10^6$ OH-radicals/cm³.

The atmospheric half-life was estimated to be 8.14 hours, the overall OH-rate constant was estimated to be $1.92*10^{-11}$ cm^{3*}molec^{-1*}sec⁻¹. The reliability of the results from the QSAR was rated Klimisch 2.

It is expected that photolytic degradation in air is no relevant pathway for removal from the environment. As it is assumed that the majority of UV-350 will be emitted indirectly via sewage treatment systems and directly via surface runoff into the aquatic compartment and considering the very low vapour pressure of UV-350 it is concluded that the substance will not evaporate at ambient temperature. This assumption is supported by the results of environmental distribution modelling (please see section 3.3.2). Therefore photolytic degradation in the atmosphere is not considered to be relevant for the PBT assessment in the light of the partition properties.

3.1.1.2.2 Phototransformation in water

Photolytic degradation of UV-350 is expected to be a relevant degradation process only in very shallow clear waters and in the first few centimetres of the water column, decreasing rapidly in the lower layers of the water column, if at all. It is expected that the environmental exposure of UV-350 occurs in the whole water column. Because of the substance's adsorption potential it will predominantly bind to suspended organic matter and sediment which is supposed to decrease the availability of the substance for photolytic degradation. Therefore photolytic degradation in the natural aquatic environment is not considered to have relevant impact on the overall persistency of UV-350.

3.1.1.2.3 Phototransformation in soil

Information from industry indicates that a small fraction of the group of phenolic benzotriazoles is used in the EU in cosmetic products. The majority of this fraction will end up in waste water and finally adsorb to sewage sludge. As the use of this sludge is a common practice in agricultural industry soil will be subject to indirect exposure. As final step the sludge will be ploughed in and therefore only negligible quantities will be available for photolytic degradation processes.

This leads to the conclusion that photolysis is not a relevant pathway for removal of UV-350 in soil.

3.1.2 Biodegradation

3.1.2.1 Biodegradation in water

3.1.2.1.1 Estimated data

To the dossier submitter's knowledge no studies exist describing the biodegradation pathway of the phenolic benzotriazoles in the environment. Therefore the pathways of all phenolic benzotriazoles in auestion were simulated with the University of Minnesota Biocatalysis/Biodegradation Prediction System (UM-PPS²). This web application is a rule- based system currently encompassing 250 microbial biotransformation rules based on over 1350 microbial catabolic reactions and about 200 biodegradation pathways. The system compares the organic functional groups of the entered molecules with its set of rules and shows all possible degradation steps. The reaction steps are color coded according to the likelihood that the respective reaction is catalysed by certain bacteria in water, soil or sediment. An overview of the system can be found in two recent publications by Ellis et al. (Ellis et al., 2008) and Gao et al (Gao et al., 2011). Please note that it is not possible to predict rate constants with this system. Also there is no defined applicability domain for this rules based system.

As the phenolic benzotriazoles are complex molecules, their degradation pathway is also quite complex. Nevertheless a comparison of the results shows similarities and patterns. To better understand the degradation processes some generalizations are helpful. For all four phenolic benzotriazoles on which dossiers were submitted three different degradation pathways are possible. The first one starts at the phenol ring of the benzotriazole moiety. While the phenol ring is degraded, this degradation pathway always ends when a triazole group is left. The second possibility is to start the degradation at the side chain in position four (para-position) to the hydroxyl group. This degradation pathway ends when the side chain is completely

² <u>http://umbbd.msi.umn.edu/predict/</u> (accessed 12.06.2012)

degraded. For the complete degradation of the phenolic benzotriazoles the third degradation pathway is the most relevant, as this one results in the degradation of the bond between the phenol ring and the benzotriazole moiety which is never directly cleaved. The UM-PPS predicts that the actual breakdown of the phenolic ring begins only when two vicinal hydroxyl groups on the phenolic ring are formed. In order to obtain the vicinal hydroxyl groups it is necessary to degrade the side chain in position six (ortho-position) first. Depending on the phenolic benzotriazole in question this encompasses many reaction steps that sometimes are not very likely (and therefore kinetically speaking slow). Of special importance in this regard is the transformation of the aliphatic methyl groups into primary alcohols. The crucial step after degradation of the side chain is reached when the two vicinal hydroxyl groups are formed. Then the carbon-carbon-bond between them is broken and therefore the phenolic ring is cleaved. The mechanism is shown in Error! Reference source not found.. In the actual degradation of the phenolic benzotriazoles all three possible degradation pathways will coexist and it is a question of the individual molecular structure of the metabolite which pathway is the kinetically most favorable. For the assessment the process was simplified by choosing the pathway that is most likely and shortest. However, it has to be noted that the rules of the UM-PPS were not explicitly derived for cleavage of phenolic rings bound to benzotriazole and therefore it is uncertain if the mechanism proposed by UM-PPS is relevant in the environment.

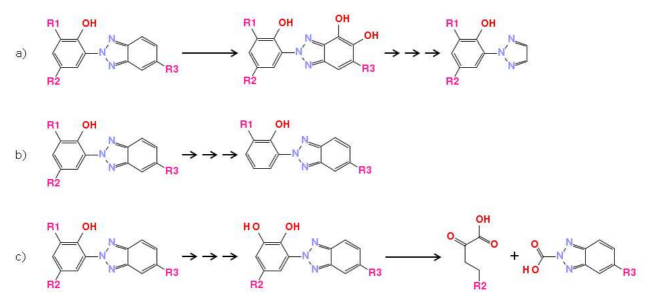


Figure 1: Proposed simplified mechanisms for the degradation of the phenolic benzotriazoles. a) Degradation of the benzotriazole moiety; b) Degradation of side chain R2; c) Degradation of side chain R1 leading to the ringcleavage of the phenolic ring R1, R2: alkyl; R3: H or Cl. Side reactions are for the sake of simplicity not considered here.

In Annex 3 an overview of the reaction pathways of all substances assessed in this chapter as predicted by the UM-PPS is given.

Since no experimental data are available on the biodegradation of UV-350 in water, a QSARcalculation with the BIOWIN module of EPISuite v4.10 (U.S. EPA, 2012) was performed. The numerical results are shown in Table 7. Details on the calculations can be found in Annex 4.

		•	
Model	QSAR result	Overall model performance	QPREF
BIOWIN2	0.1329 (does not biodegrade fast)	Reliable with Restrictions (Klimisch 2)	Annex 4.4
BIOWIN6	0.012 (does not biodegrade fast)	Reliable with Restrictions (Klimisch 2)	Annex 4.4
BIOWIN3	2.2538 (weeks to months)	Reliable with Restrictions (Klimisch 2)	Annex 4.4

Table 7: Results of BIOWIN predictions on UV-350 (reliability rated Klimisch 2)

The results of the prediction indicate that UV-350 is a borderline case for meeting the screening criteria of the ECHA Guideline R.11 for persistence. In the light of the available experimental results of similar structures (as UV-320, UV-327 and UV-350), the simulation of the prediction pathway and the discrepancies of the BIOWIN fragment approach considering the triazole group (discussed in more detail in Annex 4) it is concluded that UV-350 would meet the experimental screening criteria.

3.1.2.1.2 Screening tests

No screening tests on the biodegradability of UV-350 are available. However, studies on the structurally related benzotriazoles UV-320, UV327 and UV-328 are available, all showing little or no degradation. The results are shown in Table 8.

Substance	Result Screening Test	Test protocol	Reliability	Reference
UV-320	Non- biodegradable, BOD = 0	OECD 301C	Klimisch 2	(NITE, 2012);
UV-327	Non- biodegradable, BOD = 0	OECD 301C	Klimisch 2	(NITE, 2012);
UV-328	not readily biodegradable (2- 8% after 28 days)	OECD 301B	Klimisch 2	(The Phenolic Benzotriazoles Association, 2001);

Table 8: Results of the Degradation Screening Tests on UV-320, UV-327 and UV	
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As UV-350 is expected to follow the same degradation pattern, it is concluded in a Read-Across approach that it is not readily biodegradable, too. This finding is discussed together with the QSAR results in more detail in Annex 4.

3.1.2.1.3 Simulation tests

No simulation tests of the four phenolic benzotriazoles in question are available to the dossier submitter. However, dissipation and degradation of the substance EC 407-000-3 (Reaction mass of branched and linear C7-C9 alkyl 3-[3-(2-H-benzotriazol-2-yl)-5-(1,1-dimethyl)-4-hydroxyphenyl]propionates) in a water-sediment study according to OECD 308 was examined (dossier on 407-000-3). This study was used for a read-across on the persistence of the four phenolic benzotriazole.

Rationale for read-across asessment:

According to REACH regulation Annex XI 1.5 (Grouping of substances and read-across approach). The aim of a read-across according to REACH is to avoid testing of every substance for every endpoint, by using data known for one substance – in this case of the environmental fate - for other, similar substances. Substance similarity may be based on three criteria:

(1) a common functional group;

(2) common precursors and/or the likelihood of common breakdown products via physical and biological processes, which result in structurally similar chemicals; or

(3) a constant pattern in the changing of the potency of the properties. This criterion is of special relevance when using a grouping approach which is not done here.

Nevertheless, all three points are met: EC 407-000-3 is a phenolic benzotriazole as the four substances that are assessed in these documents. It is substituted in the positions four and six of the phenol ring just like the four substances in question. Both substitution groups are alkyl chains. Position six is substituted with a tert-butyl-group which is also present in UV-320, and UV-327. In UV-328 there is a tert-pentyl-group, the next higher homologue of a tert-butyl-group in this position. In case of UV-350 a sec-butyl-group is in position six, which is a structural isomer of a tert-butyl-group. Position four of the substances UV-320, UV-327 and UV-350 is again substituted with a tert-butyl-group, while it is substituted by a tert-pentyl-group in case of UV-328. EC 407-000-3 is substituted in position four of the phenolic ring with a propionic ester. The difference between UV-320 and UV-327 lies in a chlorine atom on the benzotriazole moiety. In summary the five substances are structurally very similar.

Not only are the substances themselves similar, but also the breakdown products are similar. The possible degradation processes for the four substances were already discussed in chapter 3.1.2.1.1. The most likely degradation pathway for EC 407-000-3 was also simulated with UM-PPS. The simplified degradation pathway is shown in Annex 3. The whole pathway follows the same pattern as observed for the four substances of interest: At first the ester is degraded in its carboxylic acid (in the following called M1, see Figure 2). Then the side chain in position four is degraded stepwise. It ends up with one of the breakdown products that are also possible for UV-320. The subsequent degradation steps are therefore the same.

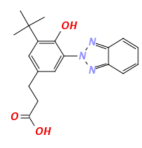


Figure 2: M1 (CAS 84268-36-0) is the first metabolite of degradation of EC 407-000-3

Based on the chemical composition of the substitution groups of the four phenolic benzotriazoles and M1 a qualitative estimation of the expected degradation times can be made:

 $\text{DegT}_{50}(\text{M1}) < \text{DegT}_{50}(\text{UV-350}) < \text{DegT}_{50}(\text{UV-328}) \approx \text{DegT}_{50}(\text{UV-320}) \approx \text{DegT}_{50}(\text{UV-327})$

The rationale can be seen in the fragment approach of Table 9.

Table 9: Fragments to be considered for qualitative assessment of degradation times.

Substance	R1	R2	R3
M1	tert-butyl	n-propionic acid	Н
UV-350	sec-butyl	tert-butyl	Н
UV-328	tert-pentyl	tert-pentyl	Н
UV-320	tert-butyl	tert-butyl	Н
UV-327	tert-butyl	tert-butyl	Cl

Therefore the degradation M1 can be regarded as a best case-scenario for the degradation half life times of the four phenolic benzotriazoles.

In conclusion the REACH criteria for applying a read-across approach are met and the degradation study of EC 407-000-3 can be used as further supporting information on degradation behaviour of the phenolic benzotriazoles.

Assessment of a a water-sediment study according to OECD 308 on EC 407-000-3 (aerobic conditions)

Test conditions are generally well described and the test was done according to GLP but validity descriptors remain unknown or even question reliability of the study, i.e. Chi² is not reported and many graphs do not sufficiently match the responding values. The report is reliable with restrictions (2 according to Klimisch).

As usual for this kind of study two systems of different organic carbon levels were employed. A river system contained low level and a pond system contained high level of organic carbon. Sampling locations of water and sediment were a pond and the river Rhine. For both systems the sampling locations were thought to not have been pre-exposed to the test substance or structural similar substances. The pond did not receive effluent discharge and this was assumed for the river Rhine, too, but as no exact sampling location was given some uncertainty remains. The test substance was radiolabelled in the benzene ring of the triazole moiety. Test systems were allowed to acclimatise for two weeks after filling. Test duration was 100 days and test temperature was 20 \pm 2 °C. As this is higher than 12°C the PBT guidance recommends to employ a temperature correction with a Q10-factor of 2.2. Please note that this factor was derived for degradation not dissipation, where it might be lower. Water sediment ratio was 3.3:1. A stock solution which consisted of test substance in aceton was stepwise diluted to give a final concentration of the test substance of 3 μ g/L. The test substance was applied dropwise onto the water surface. Water and sediment were separated and analysed at each sampling point. Two traps were employed for volatile substances. On six occasions samples were taken and analysed. Analysis was done by TLC, HPLC and LSC and recovery rate was 98.7 % (96.2-101.2 %) for the river system and 99.9 % (97.6-101.9 %) for the pond system (see Error! Reference source not found. and Error! Reference source not found.).

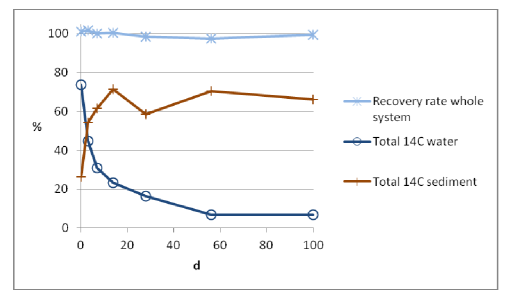


Figure 3: Recovery rate and distribution of total radioactivity in the pond system under aerobic conditions

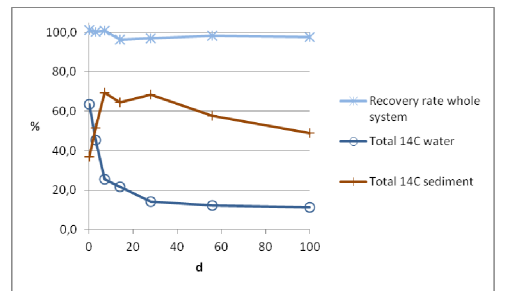


Figure 4: Recovery rate and distribution of total radioactivity in the river system under aerobic conditions

In both systems mineralisation was negligible with 1.2 or 1.3 % and the parent steadily declined to 3 or 4 % at day 100 in both systems (see Figure 5 and **Error! Reference source not found.**). The steady decline suggests cometabolic degradation processes or abiotic degradation or dissipation processes. In neither system volatile substances were detected. One metabolite (M1, CAS 84268-36-0) was identified, only. Thus, a metabolic pathway could not be substantiated although it is clear that some degradation occurred resulting in formation of the metabolite M1.

M1 **Error! Reference source not found.Error! Reference source not found.** is the respective carboxylic acid of EC 407-000-3. It was detected as the main metabolite in quantities exceeding 10 % of the applied radioactivity by far and was found as well in the water as in the sediment phase. Twelve other metabolites were detected, but not identified. Three metabolites reached amounts of 5 to 8 % each in the total system at day 100.

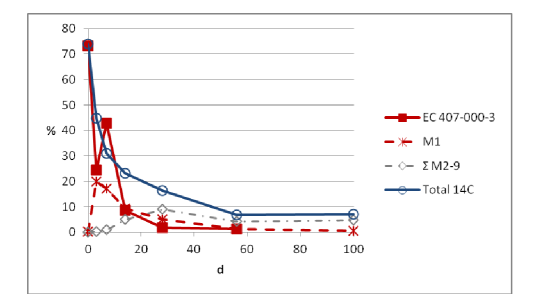


Figure 5: Parent, metabolites and total radioactivity in the water phase of the pond system under aerobic conditions

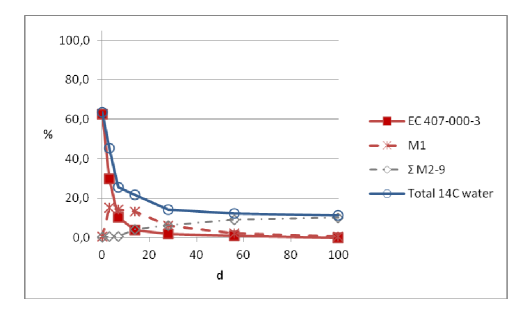


Figure 6: Parent, metabolites and total radioactivity in the water phase of the river system under aerobic conditions

The lack of mineralisation and the missing identification of further metabolites do not allow for differentiation of degradation and mere dissipation processes which contributed to the overall dissipation of M1. With no further metabolites identified adsorption and desorption of metabolites also remain unknown. Dissipation may have been caused by mere adsorption. Another aspect that hampers differentiation is the relatively high level of non extractable residues (NER), because it remains unknown to which extent parent or metabolites contributed to NER formation (see **Error! Reference source not found.** and **Error! Reference source not found.**).

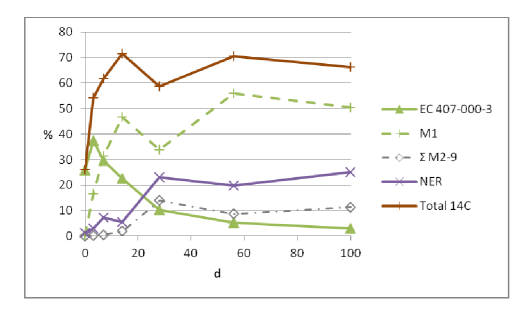


Figure 7: NER, parent, metabolites and total radioactivity in the sediment phase of the pond system under aerobic conditions

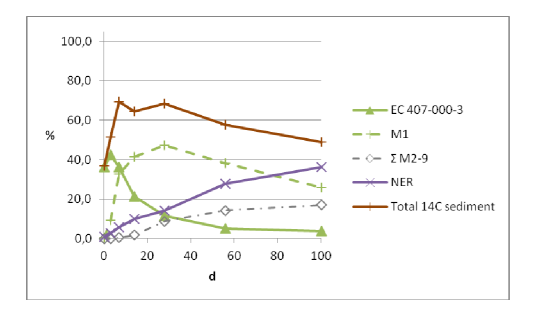


Figure 8: NER, parent, metabolites and total radioactivity in the sediment phase of the river system under aerobic conditions

In the sediment phase the trend for M1 was similar in both systems up to day 14, afterwards it differed. After reaching a maximum a clear decrease was observed in the river system, whereas only a slight decrease was observed in the pond system. In both systems the sediment values of M1 were already high at day 7 with 33 or 31 % of applied radioactivity and reached a similar high value on day 14 with 41 or 47 % (river or pond system). In the river system a maximum of approximately 47 % was reached at day 28 which finally decreased to 26 % at day 100. In the pond system an already high value of approximately 47 % on day 14 was followed by 34 % at day 28, reached a maximum of 56 % at day 56 and afterwards dropped only slightly to 50 % at test end on day 100.

In the following an attempt is made to interpret the reported concentration of M1 but it should

be kept in mind that the test was not designed to follow specifically degradation of M1. Consequently data are limited and interpretation is difficult. The major uncertainty lies in the formation of further M1 from EC 407-000-3. When doing a graphical estimation of DT_{50} this leads to an overestimation of it that will depend on the amount of parent left at this point.

In both systems M1 showed similar trends in the water phase. The maximum was reached at t day 3 (15-20% overall concentration). At day 28 the concentration had dropped below 10%, i.e. half of the concentration of the maximum at day 3. An approximate DT_{50} of 25 days results. Applying a temperature correction to this leads to a DT_{50} of 55 days. According to Annex XIII a $DT_{50} > 40$ days would show M1 to be persistent in water provided that DT_{50} would have been a $DegT_{50}$. However, in this case dissipation from the water phase to the sediment is of major importance and is very likely the the overall degradation is longer than dissipation alone, probably above 60 days.

Table 7 and Table 8 present the decline of M1 in the respective system taking the maximum value of M1 and the time at which maximum occurred as basis (see **Error! Reference source not found.** and **Error! Reference source not found.**):

Table 1: Decline of M1 for sediment and whole system concentration in the river system (low org. C)

Sediment		Whole system	
Time in d	Decline in %	Time in d	Decline in %
0	0	0	0
28	20	14	2
72	46	42	27
		86	52

Table 2: Decline of M1 for sediment and whole system concentration in the pond system (high org. C)

Sediment		Whole system		
Time in d	Decline in %	Time in d	Decline in %	
0	0	0	0	
44	10	44	11	

In the following an attempt is made to overcome the problem of a DisT_{50} probably containing degradation as well as dissipation or partitioning processes by deduction of a DegT_{50} from the specified DisT_{50} for the purpose of comparing data with trigger values.

As stated above it is not possible to differentiate between degradation and mere dissipation processes, because of missing information on real degradation and the unknown identities of the further metabolites and thus the DisT_{50} of M1 for the sediment phase represents all processes. Another aspect that hampers differentiation is the relatively high level of non extractable residues (NER), because it remains unknown to which extent parent or metabolites contributed to NER formation. NER reached 36 % in the river system and 25 % in the pond system. They were mainly bound to the humic fraction and humic acids and to a lesser part to fulvic acids. Phenolic benzotriazoles have a high log K_{OC}. Therefore they have a high tendency to adsorb.

Though data are insufficient for a detailled kinetic modelling it is possible to draw the following conclusions: $DisT_{50}$ of M1 was approximately 72 days in river system without applying temperature correction and 158 days when applying it (see Table 1 and **Error! Reference**)

source not found.). This slightly below the trigger $DT_{50} < 180$ days.

But as degradation shall be compared with the trigger value, these dissipation data are generally improper for comparison purposes. It can be stated though, that $DegT_{50}$ of M1 will be longer than 72 to 158 days because degradation is only one of all the processes which contribute to dissipation.

Some further aspects should be considered which contribute to the overall picture. In the pond system only 11 % dissipation of M1 was reached within 44 days. It is impossible to derive a DT_{50} for the pond system, not even a $DisT_{50}$. It may only be stated that $DisT_{50} > 44$ days in pond system (therefore longer than 97 days when applying temperature correction). Nevertheless, a comparison with the river data (see Table 7 and Table 8) shows that dissipation in the pond system in 44 days is only about half of the dissipation measured within the river system in 28 days which means dissipation was much slower in the pond system than in the river system.

Although it is not possible to extrapolate far beyond the available time frame the pond system data show that dissipation may be very slow depending on the conditions given.

Systems with high organic content generally should be more biologically active. They also have more potential binding sites for adsorption. The latter is thought to have been the case and would explain the different dissipation half-lives between the low and the high organic content systems.

In case of unclear contribution of partition processes to dissipation and if dissipation of the substance in question mainly takes place in sediment, the whole system should be considered, too (see **Error! Reference source not found.** and **Error! Reference source not found.**). Assessing the whole system ensures that mere adsorption will not have a decisive influence on a DT_{50} because adsorbed substance will show up in sediment and thus not dissipate in whole system .

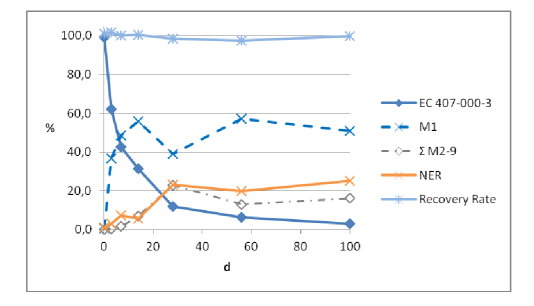


Figure 9: NER, parent, metabolites and total radioactivity in the whole system of the pond system under aerobic conditions

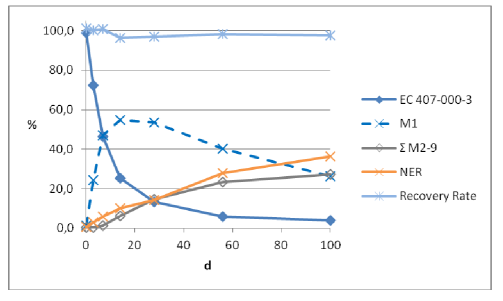


Figure 10: NER, parent, metabolites and total radioactivity in the whole system of the river system under aerobic conditions

The total occurrence of M1 (whole system) is mainly affected by M1 enrichment in sediment and consequently matches the course in sediment quite closely. Most important is the following lack of decline in the pond system (see Figure 7 to **Error! Reference source not found.**).

In both systems the whole system values of M1 were already high at day 3, increased further and reached a similar high value on day 14. In the river system a maximum of approximately 55 % was reached at day 14 which only slightly decreased until day 28 but finally decreased to 26 % at day 100. In the pond system a near maximum of 56 % was reached at day 14, dropped afterwards to 39 % and raised again reaching finally a maximum of 57 % at day 56. It only decreased slightly to 51 % at day 100. The reason remains unclear for the decline to 39 % at day 28 in the pond system. Given the overall trend it may have been an outlier possibly caused by problems in the extraction process. No such outlier was observed in the river system.

DisT₅₀ of M1 in the whole system was approximately 86 days in river system and more than 44

days in pond system. After applying temperature correction the DisT50 of the whole river system is 189 days and for the pond longer than 97 days. As degradation shall be compared with the trigger values these dissipation data are improper.

Some further aspects should be considered which contribute to the overall picture. In the pond system only 11 % dissipation was reached in 44 days (see Table 8). A comparison with the river data (see Table 7) within this time frame shows that this is only about half of the dissipation measured in the river system, i.e. dissipation was much slower in the pond system than in the river system. Moreover, dissipation may have been even much slower than this. In pond system 56 % at day 14 was observed which is as nearly as high as the maximum of 57 % at day 56 (see **Error! Reference source not found.**). Though the reported value is slightly lower it may also have been the same at both time points if one considers measuring inaccuracy. In this case 11 % of M1 would have been dissipated in 86 days.

Even though a detailed kinetic modeling is not possible a simple worst-case kinetic estimation is possible. This estimation model considers two processes, the production of M1 from the degradation of EC 407-000-3, and the further degradation/dissipation of M1, with two complementary calculations:

- A. Primary dissipation of M1 (accounting the formation from the parent)
- B. Primary and secondary degradation of M1 (dissipation of M1 and identified metabolites): the ultimate degradation time of M1 will be clearly higher than this value.

The basic assumptions are:

- A. EC 407-000-3 is degraded to M1, a manual fitting to the actual data is done.
- B. M1 is further degraded to M2-M9. This assumption is in line with the radiolabelling of the molecule, the structure, and the finding in the different systems.
- C. The modelling focuses on the last part of the experiment (the most relevant), and assumes first order kinetics for M1 allowing estimations of half-life (DT50=Ln(2)/dissipation rate)

The results for this estimation model can be inserted into **Error! Reference source not found.** and **Error! Reference source not found.** and are shown in Figure 11 and Figure 12.

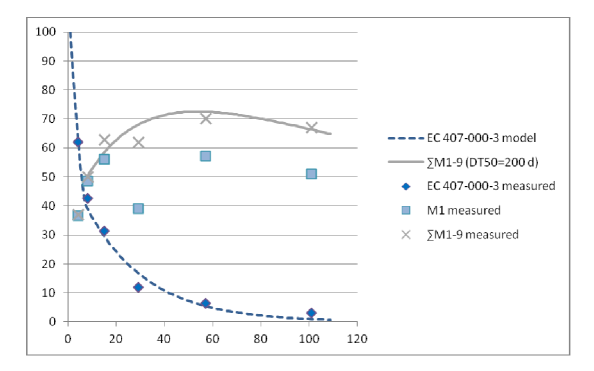


Figure 11: Parent and metabolites in the whole system of the pond system under aerobic conditions, estimation for apparent dissipation of all metabolites added (DT50 = 200 d).

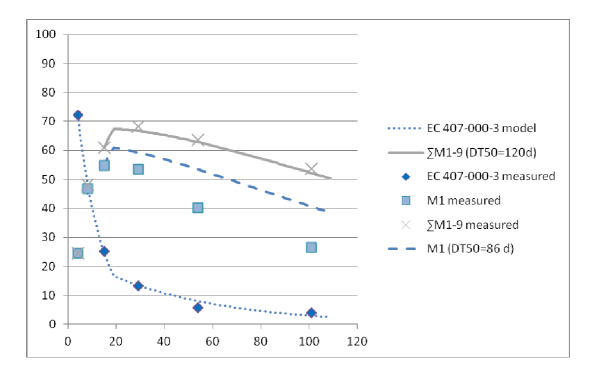


Figure 12: Parent and metabolites in the whole system of the river system under aerobic conditions, estimation for apparent dissipation of all metabolites added (DT50 = 120 d).

For both systems it is possible to model the apparent primary and secondary degradation of M1 (Σ M1-9). The estimation for the pond would lead to a DT₅₀ of 200 days and for the river to a DT₅₀ of 120 days. The ultimate degradation time of M1 will clearly be higher than this value. Please note also that in Figure 12 the estimation for M1 with a DT50 of 86 days is shown which overestimates the concentrations (see above).

Assessment of a a water-sediment study according to OECD 308 on EC 407-000-3 (anaerobic conditions)

A further test according to OECD 308 on degradation of EC 407-000-3 in water and sediment under anaerobic conditions was reported in the dossier on 407-000-3. Sediment was taken from an organic rich pond. In contrast to the aerobic test only small amounts of NER were found. With the exception of M1 all metabolites formed in small quantities, only.

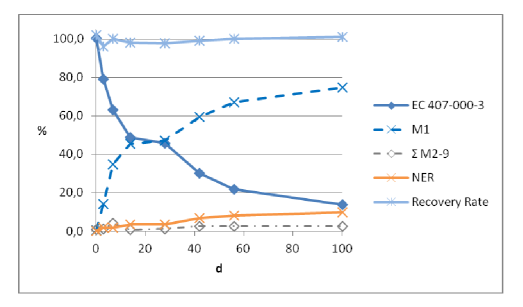


Figure 13: NER, parent, metabolites and total radioactivity in the whole system of a pond system under anaerobic conditions

M1 reached 75 % in the whole system at day 100, 65 % were located in the sediment. Up to day 14 when the maximum of 32 % was reached the majority of M1 was found in the water phase. Afterwards the concentration decreased to 10 %. In the sediment phase concentration increased to the maximum of 65 % at test end (see **Error! Reference source not found.**).

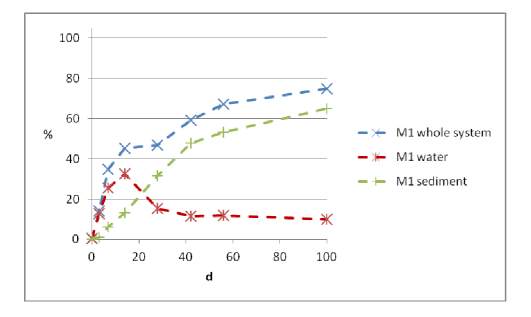


Figure 14: Main metabolite M1 in a pond system under anaerobic conditions

While EC 407-000-3 dissipated quickly its main metabolite M1 continuously built up throughout the test (see **Error! Reference source not found.**). The test was also conducted at 20°C, therefore temperature correction would have to be employed if a $DegT_{50}$ would be calculated.

As for the aerobic systems, also for this system the degradation half life was estimated by doing a worst-case model calculation. This was based on the assumptions that all of EC 407-000-3 is degraded into M1 and this would degrade by first-order kinetics. The concentrations were calculated assuming a degradation half-life of 180 days (i.e. the vP-criterion in sediments) but not regarding temperature correction. The result is shown in Figure 15.

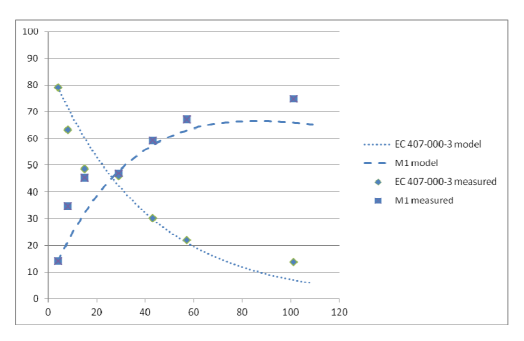


Figure 15: Model calculation on the degradation of M1 in the anaerobic system assuming a degradation half-life of 180 days (data without applying temperature correction).

The comparison of the model calculation with measured data shows that even if a DT_{50} of 180 days is assumed, the actual measured concentration of M1 is underestimated. Thus measured data indicate that at 20°C the degradation half-life is longer than 180 days.

Overall the study was not performed to determine half-lifes for the meatbolites of EC 407-000-3 and furthermore it has to be recognized that the physico-chemical properties of EC 407-000-3 and its metabolites complicate the derivation of degradation half-lifes. Nevertheless, even based on best case assumptions (i.e. favorable for degradation) it was shown that M1 will have a degradation half life of > 180 days under anaerobic conditions in sediment and in the study up to day 100 hardly any metabolites were formed and mineralization was neglectable. It is more difficult to assess the aerobic studies as under these conditions the fraction of nonextractable residues almost reaches 40% at the end of the test. Only a dissipation half life for the aerobic river system could be calclulated which was 86 days considering the whole system and 72 days considering the sediment. After applying the temperature correction this dissipation values will already be near or above the value for very perstistent substances Taking into consideration the high NER percentage and the likelyhood that dissipation does include dissipation to NER it is very likely that the overall $DeqT_{50}$ will be higher. Simple kinetic modeling for the three systems at 20°C shows that the DT50 of the river system will be around 120 days, around 200 days for the aerobic pond and higher than 180 days for the anaerobic pond.

Even it can not be proven that the vP trigger is reached it has to be considered that the data show that it has to be considered that M1 does not rapidely degrade and will therefore be distributed into the anaerobic part of the sediment, where it will be even more persistent.

As M1 is a best case read-across example for the phenolic benzotriazoles in question, it was concluded that they will be also very persistent.

3.1.2.2 Biodegradation in sediments

Data from a Water-Sediment Test according to OECD 308 on the substance EC 407-000-3 (Dossier on EC 407-000-3) show that sediment is a sink for the metabolite M1 (*cf.* 3.1.2.1.3). It is not possible to derive a $DegT_{50}$ but only a $DisT_{50}$ which is improper for comparison with the trigger values. This tentative $DisT_{50}$ is > 44 days or approximately 72 days depending on organic carbon content of the system for aerobic conditions. Under anaerobic conditions M1 is very persistent because it continuously built up throughout the test.

3.1.2.3 Biodegradation in soil

No data available.

3.1.2.4 Summary and discussion on biodegradation

Although there are no simulation tests on UV-328 itself, the results of the screening test as well as the result of simulation of these tests indicate a very low potential for biodegradation. The assumed degradation pathway is similar for all phenolic benzotriazoles and starts with a degradation of the side chains that are in ortho-position to the hydroxyl group of the phenolic ring. There is a simulation study on EC 407-000-3 which also gives information on a metabolite having a similar structure to the phenolic benzotriazoles in question. As it can be assumed that this phenolic benzotriazole will also be biodegraded according to the same mechanism and as it is structurally very similar to the four phenolic benzotriazoles the results of this substance can be used as an argument for read-across. As from a qualitative point of view M1 will degrade faster, this is a read-across on a best case example. Though it is impossible to compare data

directly with the trigger values data give enough information to conclude that degradation will be very slow under predominant aerobic conditions in environment (based on model calculations longer than 120 days in the river system and longer than 200 days in the pond system). It is believed that adsorptive substances will reach anaerobic zones in sediment sooner or later. Therefore anaerobic biodegradation is of special interest for these substances. M1 was constantly built up under anaerobic conditions and was hardly degraded at all. The degradation process of the parent obviously stopped at this stage. A model calculation using worst case assumptions showed that the degradation half life is > 180 days. UV-350, which has a sec-butyl-group as side chain in ortho-position that is approximately not quite as hard to degrade as a tert-butyl-group, will accordingly have a degradation half-life time that is a bit smaller, but nevertheless comparable. This is supported by the simulated degradation pathway.

3.1.3 Monitoring studies

For UV-327 and UV-328 four studies are available which investigated the distribution of UVthem in sediments in a highly contaminated area (Narragansset Bay, Rhode Island, USA). Taken together, the information can be used to find some hints about the degradation potential of the phenolic benzotriazoles in sediments.

UV-327 and UV-328 were historically produced in an industrial plant at the Pawtuxet river which flows into the brackish Providence River and consequently the Narragansset Bay (Reddy et al. 2000, Junclaus et al. 1980 and Lopez-Avila and Hites, 1980). Production of UV-327 was reported between 1963 and 1972, while UV-328 was produced from 1970 to 1985 (Hartmann et al. 2005, Lopez-Avila and Hites, 1980).

Two studies provide information about the sediment concentration during the production phase:

Jungclaus et al. (Jungclaus et al., 1978) analyzed industrial WWTP effluent and receiving waters and sediments from that American specialty chemicals manufacturing plant producing organic compounds and running a badly performing WWTP. 16 water samples and 19 sediment samples (located at different sites at the Pawtuxet river including the Pawtuxet cove) were taken in 1975 and 1976 and the compounds contained were identified, beside others UV-327 and UV-328. River water and sediments were collected in Providence River and its tributary Pawtuxet River (Pruell et al., 1984). UV-328 was detected in industrial WWTP effluent (0.55 – 4.7 ppm), in river water (7 – 85 ppb) and in sediments (1-100 ppm). UV-327 was detected only in sediment, with concentrations of 2 – 300 ppm.

(Lopez-Avila and Hites, 1980) investigated the same specialty chemicals manufacturing plant located on the Pawtuxet River. Eight sediment cores were taken at three locations in the Pawtuxet River. The sites were selected for an abundance of fine-grained material. Further sediment cores were taken at four locations in the Pawtuxet Cove and 13 locations in the Providence River and Narragansett Bay. The core concentrations of the compounds in the sediment have been condensed into a single number. However, the authors feel the values given are representative of the sediment concentrations. Concentrations decrease both with depth in the sediment and with increase in distance from the discharge.

	Pawtuxet River			Pawtuxet	Providence	River	
	near plant	mid river	near dam	Cove	near	far	bay
UV-327	300	400	20	80	20	2	0.5
UV-328	300	300	70	100	10	5	0.6

Table 10: Concentrations of phenolic benzotriazoles in sediment cores (ppm)

In summary both studies show that sediment concentrations were in the range of 2-300 ppm (UV 327), and 1-300 ppm (UV 328) in the Pawtuxet river and the providence river while concentrations in the Narragansett Bay was lower (0.5 ppm for UV 327 and 0.6 ppm for UV 328) during the actual production of these compounds.

Two further studies provide some evidence about the concentration of the compounds years after the production has ceased:

Reddy et al. (Reddy et al., 2000) examined the free and bound fractions of different substituted benzotriazoles in sediment cores from the Pawtuxet River and Narragansett Bay in the U.S. The Pawtuxet River sediment core was collected in 1989 and sectioned at 2-3 cm intervals. Eleven sections from 0-2 cm to 50-52 cm were analyzed. The sedimentation rates in this section of the river are 2-3 cm/year. The redox discontinuity, determined visually, was in the top 2 cm of the core. The Narragansett Bay core was collected in 1997. Six sections from the top 13 cm of the core were analyzed. The sediments in this area become anoxic within a few millimeters of the surface and have a sedimentation rate of about 0.3 cm/year. The method detection limit was ca. 20 ng/g for each (free and bound) fraction.

In the Pawtuxet River core no bound benzotriazoles were detected. UV-327 was most abundant: the highest concentration was ca. 5 mg/g and it was observed down to 50-52 cm. The graph shows a varying concentration in the first 20 cm and a constant decrease with depth starting at 20-22 cm. Taking into account a sedimentation rate of 2-3 cm/year for this site, a depth of 20 cm means that exposure was 7-10 year before the actual measurement. If it is assumed that exposure was constant during the years, the decrease in the UV-327 concentration between 20 and 50 cm depth should reflect the degradation rate of UV 327. As a very rough estimate concentration decrease in depth can be compared to a decrease which would be expected assuming a DegT50 of 180 days (see Table 11, assumption: 2.5 cm depth reflects 1 year)

Depth [cm]	approximate measured c	expected c assuming a DegT ₅₀ of 180 d [mg/kg]
20	10 ⁵ ng/g (100 mg/kg)	
25	4 x 10 ³ ng/g (4 mg/kg)	25
30	6 x 10 ² ng/g (0.6 mg/kg)	1
40	3 x 10 ² ng/g (0.3 mg/kg)	0.15
52	10 ² ng/g (0.1 mg/kg)	0.075

Table 11: Concentration profile of UV 327 based on a graphical evaluation from Reddy et al. (2000) and expected concentration based on a $DegT_{50}$ of 180 d at the different depths

Although this is a very rough estimation for which uncertainties need to be taken into account it supports a very slow degradation of UV-327.

In addition the study, as well as a second study by Hartmann et al (2005) can be used to compare actual concentration with historical data which also may provide some information about the degradation time (see below)

Hartmann et al. (2005) took sediment cores at three locations in Narragansett Bay in 1997 (Apponaug Cove, Seekonk River, Quonset Point). The cores were analyzed for several contaminants including UV-327 and UV-328. Two of the cores were split into 2 cm sections, and the third core (Quonset Point) was split into 10 cm sections.

The concentrations of UV-327 and and UV-328 at the different depth are summarized in table 45.

Quonset Po	oint core		Apponaug	Cove core	Seekonk River core		
depth [cm]	UV-327 [ng/g dw]	UV-328 [ng/g dw]	depth [cm]	UV-327 [ng/g dw]	UV-328 [ng/g dw]	UV-327 [ng/g dw]	UV-328 [ng/g dw]
0 – 2	ca. 40	ca. 160	0 - 2	ca. 130	ca. 270	ca. 30	ca. 120
0 - 10	ca. 60	ca. 260	2 - 4	ca. 30	ca. 80	ca. 20	ca. 70
10 - 20	ca. 80	ca. 360	6 - 8	ca. 50	ca. 140	ca. 30	ca. 140
20 - 30	ca. 100	ca. 840	10 - 12	ca. 70	ca. 120	-	-
30 - 40	ca. 130	ca. 1100	12 - 14	-	-	ca. 5	ca. 20
40 - 50	ca. 690	ca. 1180	20 - 22	n.d.	n.s.	n.d.	n.d.
50 - 60	ca. 480	ca. 40	30 - 32	n.d.	n.d.	-	-
60 - 70	n.d.	n.d.	38 - 40	-	-	n.d.	n.d.
80 - 90	n.d.	n.d.	40 - 42	n.d.	n.d.	-	-
100 - 110	n.d.	n.d.	48 - 50	-	-	n.d.	n.d.
119 - 129	n.d.	n.d.					

Table 12: Concentrations of phenolic benzotriazoles in sediment cores from Narragansett Bay (concentrations taken from a graph)

n.d. = not detected

not measured

Taking into account the specific sedimentation rate at each site, it is possible to identify the layer which probably represents exposure during active production of UV-327 and UV-328. This might be used – as a very rough estimate to compare concentrations with historical concentrations during production in order to get an idea about whether or not degradation occurred. Unfortunately historical data are not available for the three sampling site and thus the comparision is highly uncertain. However, as a second type of information the data can be used to calculate how high concentrations can be done based on the results of Reddy et al. (Reddy et al., 2000) (see above) The results of these calculations are summarized in Table 13.

Study	Detecti on limit [ng/g]	Site	Year of collectio n	Sedime ntation rate [cm]	Layer assumed to reflect production period [cm]	c at that layer	estimated c during production (if DegT ₅₀ =180d)	probably not at the
UV 327 (production	period 196	53 -1972					I	
Reddy et al., 2000	20	Pawtuxet River	1989	2-3	34 -69	10 ² ng/g (0.1 ppm) (At 52 cm)	13107 ppm	20 – 300 ppm (Junghans et al, Pawtuxet river)
Hartmann et al	10	Quonset Point (Narragan sett Bay)	1997	2	54 - 68	~ 500 ng/g (0.5 ppm) (at 50 - 60 cm)	Not possible	0.5 ppm ((Lopez-Avila and Hites, 1980, Narragansett Bay)
Hartmann et al	10	Apponaug Cove (Narragan sett Bay)	1997	0.5 – 0.85	14 - 29	~ 70 ng/g (0.07 ppm) (at 10 -12 cm)	Not possible	0.5 ppm ((Lopez-Avila and Hites, 1980, Narragansett Bay)
UV-328 (production		70 – 1985)						
Hartmann et al	10	Quonset Point	1997	2	24 – 54	~ 40 ng/g (0.04 ppm) (at 50 - 60 cm)	9175 ppm	0.6 ppm ((Lopez-Avila and Hites, 1980, Narragansett Bay)
Hartmann et al	10	Apponaug Cove	1997	0.5 – 0.85	6 - 23	~ 130 ng/g (0.13 ppm) (at 10 -12 cm)	17039 ppm	0.6 ppm ((Lopez-Avila and Hites, 1980, Narragansett Bay)

Table 13: comparison of estimated historical concentrations based on a DegT50 of 180d and historical concentrations from literature

Although these data are highly uncertain they show that the assumption of a degradation half life of 180 days leads to unrealistic high starting concentrations and therefore this provides further support for the assumption that degradation of UV-327 and UV-328 in sediments is expected to be very slow, with a degradation half time above 180 days.

There are only two monitoring studies available on UV 350. Only in one of the studies environmental matrices were analyzed, the other study relates to WWTP sludge. No conclusions can be drawn from these data.

3.1.4 Summary and discussion on degradation

Biodegradation is expected to be the most relevant pathway for degradation of UV-350, if there is degradation. The overall evidence presented in chapter 3.1.2 in combination with the high-potential for adsorption on soil and suspended organic particles indicate in a Weight-of-Evidence Approach that UV-350 will be persistent in the environment. This is based on the following findings:

The QSAR-calculations on UV-350 indicate that it will be at least a borderline case for meeting the screening criterion for persistence. Combined with the available experimental data of ready biodegradation tests on UV-320, UV-327 and UV-328 it plausible, that UV-350 will also have a loew potential for biodegradation in these tests.

The simulation study on EC 407-000-3 is used for a read-across-assessment on a best case example, namely the first metabolite of the substance which is its carboxylic acid. The study has several shortcomings for using in the assessment, most of all that it was not conducted to assess the fate of the metabolites. Also the study lasted only for 100 days, therefore, the results have to be extrapolated to compare them with the relevant trigger values for assessing persistence in sediment. Finally, the physico-chemical properties of the substance complicate matters as considerable amounts of substances are bound in non-extractable residues especially in the aerobic systems. Nevertheless it is possible to derive important information on the persistence of phenolic benzotriazoles from this test. Though it is not possible to derive half-lives for degradation from the system, for the river system where the largest amount of metabolites are found a temperature corrected $DisT_{50}$ of 189 days is found. In the anaerobic pond system the metabolite M1 is formed up to the end of the test and only small amounts of other metabolites are detected. Starting from this data it is possible to model the expected concentrations of M1 considering a case where all of EC 407-000-3 will form M1 and this degrades by a first order kinetics. The model calculation for the data at 20° C shows that the actual half-life has to be > 180 days.

The four monitoring studies on UV-327 and UV-328 from Rhode Island show how phenolic benzotriazoles will persist in the environment. In these studies the concentrations that where found when the two substances were produced are given as well as the concentrations that were found up to 25 years later. It is not possible to derive reliable $DegT_{50}$ from these studies. Also caution is needed when comparing

the data as for each study different sampling sites and methods were employed. Also, an exact description of the samples is missing (e.g. oxygen content, further contaminents, etc.). Nevertheless, from the available on one study it is possible to successfully semiquantitatevly model the concentration curve assuming slow degradation. Also, as we have some information on sampling sites and the respective sedimentation rates it is possible to assign the concentration found years after production has ceased to certain production years. With this information we can very roughly estimate the starting concentration if we assume a certain half-life. When assuming a DegT₅₀ of 180 days the resulting concentrations are completely unrealistic high. Therefore the DegT₅₀ of UV-327 and UV-328 has to be > 180 days. In a read-across assessment it is plausible that the DegT₅₀ of UV-350 will be of comparable length.

Further information supports the most important findings above:

With help of UM PPS system the three complex degradation pathways for the phenolic benzotriazoles were simulated. Only one of the three will lead to complete mineralization. As the pathways are similar for the four substances as well as for the substances where a read-across is employed it is possible to generalize the individual findings and rationalize the similarities from a mechanistic point of view. The UM PPS has the drawback that it is not possible to employ quantitative kinetic models and there are also no studies known to us that prove them to be correct. Nevertheless, from the chemical point of view they predict very plausible pathways and address all possibilities

3.2 Environmental distribution

3.2.1 Adsorption/desorption

As there is no registration dossier available QSAR-based calculations were performed to estimate the adsorption behaviour to soil or suspended organic matter for this substance. Details of the prediction can be found in Annex 5, the default input parameters were used.

Model	QSAR result	Overall model performance	QPREF	
EPISuite 4.1 KOW-	K _{oc} (L/kg): 4.52 10 ⁴	Reliable with Restrictions	Annex 5.4	
method	Log K _{oc} : 4.66	(Klimisch 2)		
EPISuite 4.1 MCI-	K _{oc} (L/kg): 1.56 10 ⁵	Reliable with Restrictions	Annex 5.4	
method	Log K _{oc} : 5.19	(Klimisch 2)		
COSMOtherm	K _{oc} (L/kg): 7.94 10 ⁴	Reliable with Restrictions	Annex 5.4	
	Log K _{oc} : 4.90	(Klimisch 2)		

Table 14: Results adsorption behaviour predictions of UV-350

The results of the estimation of the adsorption behaviour lead to the conclusion that UV-350 will strongly adsorb to organic material.

3.2.2 Volatilisation

The tendency for volatilization from the water phase was estimated by calculation of the Henry constant. Using the physical-chemical substance properties from Table 6 and a water solubility of 0.1395 mg/l (QSAR estimation from log K_{OW} with the EPISuite module WSKOW v1.41, please note that the log K_{OW} was estimated with EPISuite as well), the calculated Henry constant³ was determined to be $1.614*10^{-3}$ Pa*m³*Mol⁻¹ indicating only little tendency for volatilization. The air-water partitioning coefficient (K_{air-water}) may be derived from the Henry's law constant and is calculated to be $6.81*10^{-7}$ m³/m³. As K_{air-water} and Henry's law constant are manually calculated from QSAR-based physical-chemical substance properties the reliability of the values is rated Klimisch 2.

The $K_{air-water}$ and Henry's law constant are very low suggesting that volatilisation is unlikely to be a significant removal mechanism for UV-350 from aquatic systems and it is unlikely that the substance will be transported very far in the atmosphere (based on its atmospheric half-life estimated to be 8.14 hours).

3.2.3 Distribution modelling

Fugacity Level III distribution modelling

When released to the environment UV-350 will be distributed to the environmental compartments in different amounts. The table below shows the result of Fugacity Level III distribution modelling using EPI Suite v4.10 with the substance properties calculated within EPI Suite . The reliability of the result from the EPI Suite calculation is rated Klimisch 2.

Table 15: Distribution according to Mackay Level III Fugacity Model (estimation with	
standard parameters as provided by EPI Suite v4.10)	

compartment	mass amount (percent)		
Air	4.64*10 ⁻⁵		
water	5.21		
soil	58.5		
sediment	36.3		

The results of the distribution modelling and physical-chemical substance properties lead to the conclusion that the overall amount of the substance will adsorb to the soil (58.5%) and the sediment (36.3%).

Distribution in waste water treatment plants

The dominant route of exposure for UV-350 is expected to be wastewater which is

³ according to equation R.16-4 from ECHA Guidance on Information requirements and Chemical Safety Assessment – Part R.16 (May 2010)

treated in sewage treatment plants. Therefore distribution modelling based on physical-chemical data from table 5 and a water solubility of 0.1395 mg/l (QSAR estimation from log K_{OW} with the EPISuite module WSKOW v1.41) has been conducted to estimate the distribution of the substance in sewage treatment plants with the help of SimpleTreat. The calculation was done assuming that the substance is not readily biodegradable (k=0/h) and the reliability was rated Klimisch 2.

Table 16: Distribution in sewage treatment plants (acc. To SimpleTreat 3.0, debugged version; 7 Feb 1997)

Summary of distribution	percent		
to air	0.0		
to water	9.2		
via primary sludge	65.8		
via surplus sludge	25.0		
Degraded	0.0		
total	100		

The results of the calculation leads to the conclusion, that when UV-350 is released into waste water the largest part of the substance will be hold back in the sewage sludge and does not enter the environment. This is in agreement with available experimental findings of Ruan et al. (Ruan et al., 2012), see Annex 7. It has to be kept in mind that the use of sludge from municipal sewage treatment plants for agricultural purposes is a common practice in many regions. Over this way the substance might be released into agricultural soil.

3.3 Bioaccumulation

To our knowledge there are no experimental log K_{ow} -values for UV-350. Therefore the value was calculated with the QSAR model KOWWIN of EPISuite 4.10 and with COSMOtherm. Details on these calculations can be found in Annex 6.

Model	QSAR result	Overall model performance	QPREF
EPISuite 4.1 KOWWIN	Log K _{ow} : 6.31	Reliable	Annex 6.3
COSMOtherm	Log K _{ow} : 7.11	Reliable	Annex 6.3

Table 17: QSAR-results for log KOW-predictions of UV-350

Based on the estimated log K_{OW} -values that are larger than 4.5, it is expected that UV-350 will bioaccumulate.

3.3.1 Aquatic bioaccumulation

UV-350 was tested in a bioconcentration study according to OECD 305 C (NITE, 2012; reliability rated Klimisch 2). Not all test conditions can be reported because the summary of the studies does not list them. Two substance concentrations were

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tested in common carp (*Cyprinus carpio*). The test duration was each time 60 days. No information on the use of a dispersant is given, but in two similar studies on UV-327 which has also a low water solubility dispersants were used. Table 18 lists the original report data amended with the BCF normalised to 5 % lipid content calculated with the average lipid content of 1.89 % (lipid content 1.47 % at start and 2.31 % at end of test).

Table 18: BCF reported and BCF lipid normalised of UV-350 (values refer to whole body wet weight basis unless no other information is provided)

Test concentration in µg/L	BCF _{reported}	BCF _{lipid} -normalised		
1.0	7700 ¹	20263		
0.1	13000 ¹	34210		

BCF values are clearly well above the vB-criterion.

Additional BCF data for skin, head, innards and edibles and depuration data was received by NITE:

Table 19: Reported tissue BCF

Test concentration in µg/L	Skin	Head	Innards	Edible
1.0	16000, 8900	17000, 11000	29000, 17000	11000, 5500
0.1	13000, 17000	18000, 25000	31000, 52000	9000, 12000

Clearance half-lives of 15 and 14 days were reported for the separate test concentrations in the order as stated above.

3.3.2 Terrestrial bioaccumulation

No data available.

3.3.3 Summary and discussion of bioaccumulation

Data show very high bioconcentration factors for both test concentrations. Hence, UV-350 clearly meets the vB-criterion.

Structural similar substances confirm the assessment. UV-320 (CAS 3846-71-7) and UV-327 (CAS 3864-99-1) have been shown to meet the vB-criterion as well. Additionally, enrichment in a top predator was observed for UV-327 and UV-328 (CAS 25973-55-1).

There are no monitoring data available on UV-350 in biota.

Table 20 gives an overview over the available data on bioconcentration on all four phenolic benzotraizoles discussed.

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Table 20: Overview of the available data on bioconcentration properties of UV-320,
UV-327, UV-328 and UV-350 (values refer to whole body wet weight basis unless no
other information is provided)

Substance	Species	BCF/BAF (lipid norm.)	с [µg/L]	Test system	Туре	References
UV-320	Cyprinus carpio	1,945*	10	OECD 305C	kinetic	(NITE, 2012)
		5,905*	1			
		12,041*	0.1			
UV-327	Cyprinus carpio	1,203	1.0	OECD 305C	steady state	(NITE, 2012)
		6,283	0.1			
		8,817	0.1			
		7,540	0.01			
	Neophocaena phocaenoides	5,946	0.012 **	Monitoring	-	(Nakata et al, 2010)
UV-328	Cyprinus carpio	1,121	0.1	OECD 305C	steady state	(NITE, 2012)
		740-2,148	0.01			
		3,681	0.01			
		1333-3309	0.8	OECD 305C	-	(Ciba, 2000)
		2738-6650	0.08			
UV-350	Cyprinus carpio	20,263	1.0	OECD 305C	steady state	(NITE, 2012)
		34,210	0.1			

* at test end

** geometric mean concentration reported by Ministry of Environment, Japan

3.4 Secondary poisoning

UV-350 is expected to enrich in top predators because accumulation through the food chain was shown for the structural similar UV-327 and UV-328. Several biomonitoring studies suggest that as well (see Annex 7).

4 Human health hazard assessment

Not relevant for the identification of this substance as SVHC in accordance with Article 57(e).

5 Environmental hazard assessment

5.1 Aquatic compartment (including sediment)

5.1.1 Toxicity data

5.1.1.1 Fish

5.1.1.1.1 Short-term toxicity to fish

Under the Toxic Substances Control Act of US EPA an acute toxicity study on *Oryzias latipes* was conducted in 2001 by Ciba-Geigy (Ciba-Geigy, 2001). According to the result summary the LC_{50} (96 h) was larger than 250 mg l⁻¹ meaning that no effect was observed up to the water solubility limit. Unfortunately only an excerpt summarizing the original study report was available for decision making. Therefore no information of any analytical follow-up of the test concentrations can be provided.

Species	Duration	LC ₅₀ (mg l ⁻	Method, conditions	Reliabilty	Reference
Oryzias latipes	96 h	>250	Japanese Industrial Standard (JIS K 0102-1998- 71.), "Testing methods for industrial waste water, Acute toxicity test with fish"	Klimisch 2	(Ciba-Geigy, 2001)

Table 21: Acute toxicity of UV-350 on fish

5.1.1.1.2 Long-term toxicity to fish

No data relevant for assessing the T-criterion can be reported.

5.1.1.2 Aquatic invertebrates

No data relevant for assessing the T-criterion can be reported.

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5.1.1.3 Algae and aquatic plants

No data relevant for assessing the T-criterion can be reported.

5.1.1.4 Sediment organisms

No data relevant for assessing the T-criterion can be reported.

5.1.1.5 Other aquatic organisms

No data relevant for assessing the T-criterion can be reported.

5.2 Terrestrial compartment

No data relevant for assessing the T-criterion can be reported.

5.3 Atmospheric compartment

No data relevant for assessing the T-criterion can be reported.

5.4 Microbiological activity in sewage treatment systems

No data relevant for assessing the T-criterion can be reported.

5.5 Non compartment specific effects relevant for the food chain (secondary poisoning)

No data relevant for assessing the T-criterion can be reported.

5.6 Toxicity test results concerning endocrine disruption relevant for the environment

As there is some discussion on endocrine disrupting properties data on this issue was compiled in Annex 8.

6 Conclusions on the SVHC Properties

6.1 **PBT**, vPvB assessment

6.1.1 Assessment of PBT/vPvB properties – comparison with the criteria of Annex XIII

6.1.1.1 Persistence

If UV-350 is degraded, biodegradation is expected to be the most relevant pathway for degradation.

There are no degradation simulation tests on UV-350 itself. Nevertheless Annex XIII allows conclusions on the persistence according to REACH Annex XIII 2. REACH Annex XIII 2 in turn allows the assessment of PBT-properties in a weight-of-evidence approach, as defined in REACH Annex XI 1.2. This means that information from several independent sources is considered to conclude on a dangerous property, in this case the persistence. While each single source of information might be regarded as insufficient to support the conclusion, the combined information is, due to its weight of evidence, regarded to be sufficient.

In case of UV-328 the weight-of-evidence-approach is based on the following important facts:

- The results of the screening test on UV-320, UV-327 and UV-328 along with the QSAR-results on UV-350 indicate a very low potential for biodegradation
- There is a simulation study on a very similar phenolic benzotriazole that available allows a read –across assessment: While the study on EC 407-000-3, a similar substance which should degrade faster, does not allow a direct comparison of data with the trigger values, it shows that even dissipation of its first metabolite is very slow (DisT₅₀ of 86 days in a river system at 20°C, 189 days when temperature corrected for 12°C). Thus degradation will be even slower. This metabolite is hardly degraded at all under anaerobic conditions and the degradation half-life for this is > 180 days even at 20°C. Considering the high potential for adsorption these conditions are expected to be of special importance and anaerobic soil and sediments are expected to be substance sinks, as the substance does not rapidly degrade.
- There is a case of several studies on deeper sediments on Rhodes Island, where UV-327 and UV-328 are found in sediments up to 25 years after the production of the substances in a nearby chemical plant has stopped. Estimations on concentration curves indicate that the DegT50 has to be larger than 180 days to explain the findings.

Furthermore there is additional data supporting this assessment:

 Once released into the environment most UV-350 will be bound to soil and sediment as the substance has a very high potential for adsorption. This was demonstrated by experimental results on sewage sludge as well as simulated log K_{OC} values.

• In the common relevant mechanism for degradation of phenolic benzotriazoles the side-chain in ortho-position is degraded. The more complex this side chain is, the longer it will take for the respective substance to be degraded. In case of UV-350 a sec-butyl group has to be degraded.

6.1.1.2 Bioaccumulation

UV-350 shows very high bioconcentration in Carp with BCF exceeding the vB trigger of 5,000 by far. This finding is in line with BCF of the other benzotriazoles UV-320 and 327. Additionally, enrichment at the top of the food chain has been proven for UV-327 and UV 328. Thus UV-350 is very bioaccumulative.

6.1.1.3 Toxicity

The available studies show that UV-350 is not acutely toxic for aquatic organisms. There is no information on the long-term toxicity of UV-350. Based on the currently available data it is concluded that UV-350 does not fulfil the T-criterion.

6.1.2 Summary and overall conclusions on the PBT, vPvB properties

According to a weight-of-Evidence argumentation UV-350 has to be considered vP and therefore also P.

Overview of the conclusions of the weight-of-evidence approach:

- QSAR calculations indicate that UV-350 is a borderline case for meeting the screening criterion for persistence. Also, the structural similar substances UV-320, UV-327 and UV-328 all show a low biodegradability in ready biodegradation tests. Therefore in a read-across it is plausible that this will be the case for UV-350 as well.
- Read-across assessment on EC 407-000-3 and its first metabolite: Very slow dissipation in aerobic systems (sediment and water) near or above the vPtrigger value based on data for the different compartments with and without temperature correction. Modelling of anaerobic system shows a DegT50 > 180 days already at 20°C. Degradation of the substances in question has to be even longer;
- For UV-327 and UV-328 there are monitoring studies available showing that the substances were found decades after environmental exposure has stopped. Model calculations indicate that these findings can only be explained if the DegT₅₀ is larger 180 days.
- Further supporting information:
 - $_{\rm O}$ Simulation of the complex degradation pathways gives a mechanistic

explanation for similarities and findings;

Thus, applying the weight-of-evidence approach the substance fulfills the P and the vP-criterion of REACH Annex XIII.

Based on a MITI-BCF study the substance fulfils the B and the vB criterion of REACH Annex XIII.

In conclusion UV-350 meets the criteria for a vPvB-substance according to Article 57 e).

6.2 CMR assessment

Not relevant for the identification of this substance as SVHC in accordance with Article 57(e).

7 References

US EPA. 2011. Estimation Programs Interface Suite[™] for Microsoft® Windows, v 4.1. United States Environmental Protection Agency, Washington, DC, USA.

Ellis LBM, Gao J, Fenner K, Wackett LP. 2008 Jun. The University of Minnesota pathway prediction system: predicting metabolic logic. Nucleic Acids Res 36:W427-W432.

Gao J, Ellis LBM, Wackett LP. 2011 Apr. The University of Minnesota Pathway Prediction System: multi-level prediction and visualization. Nucleic Acids Res 39(Web Server Issue):W406-W411.

The Phenolic Benzotriazoles Association. 2001. High Production Volume (HPV) Challenge Program - Data Summary and Test Plan for Phenolic Benzotriazoles. 1-139.

Liu YS, Ying GG, Shareef A, Kookana RS. 2011 Jula. Biodegradation of three selected benzotriazoles under aerobic and anaerobic conditions. Water Res 45:5005-5014.

Ruan T, Liu R, Fu Q, Wang T, Wang Y, Song S, Wang P, Teng M, Jiang G. 2012 Jan. Concentrations and Composition Profiles of Benzotriazole UV Stabilizers in Municipal Sewage Sludge in China. Enivronmental Science and Technology 46:2071-2079.

NITE. 2012. Chemical Risk Information Platform (CHRIP).

Ciba-Geigy. 2001. Bioconcentration test of TINUVIN 343 in carp - final report. 1-100.

Hansch, C. et al: Exploring QSAR Vol 2: Hydrophobic, Electronic, and Steric Constants (1995)

Miller D, Wheals BB, Beresford N, Sumpter JP. 2001 Feb. Estrogenic activity of phenolic additives determined by an in vitro yeast bioassay. Environmental Health Perspectives 109(2):133-138.

Kawamura Y, Ogawa Y, Nishimura T, Kikuchi Y, Nishikawa J, Nishihara T, Tanamoto K. 2003. Estrogenic Activities of UV Stabilizers Used in Food Contact Plastics and Benzophenone Derivatives Tested by the Yeast Two-Hybrid Assay. J Health Sci 49(2):205-212.

Kunz PY, Galicia HF, Fent K. 2006 Jan. Comparison of In Vitro and In Vivo Estrogenic Activity of UV Filters in Fish. Toxicol Sci 90(2):349-361.

National Institute of Environmental Health Sciences. 2011. Chemical Information Review Document for Phenolic Benzotriazoles - Supporting Nomination for Toxicological Evaluation by the National Toxicology Program . 1-149.

Rodríguez Pereiro I, Casado Agrelo J. 2012. Benzotriazole UV Stabilizers in Soil and Suspended Particulate Matter Samples.

Brorström-Lundén E, Remberger M, Kaj L, Hansson K, Andersson H, Haglund P, Andersson R, Liljelind P, Grabic R. 2011. Screening of benzothiazoles, benzenediamines, dicyclohexylamine and benzotriazoles 2009. 1-64.

Carpinteiro I, AbuÃ-n B, RodrÃ-guez I, Cela R, Ramil M. 2010a. Headspace solidphase microextraction followed by gas chromatography tandem mass spectrometry for the sensitive determination of benzotriazole UV stabilizers in water samples. Analytical and Bioanalytical Chemistry 397(2):829-839.

Carpinteiro I, AbuÃ-n B, RodrÃ-guez I, Ramil M, Cela R. 2010b. Pressurized solvent extraction followed by gas chromatography tandem mass spectrometry for the determination of benzotriazole light stabilizers in indoor dust. Journal of Chromatography A 1217(24):3729-3735.

Carpinteiro I, Ramil M, RodrÃ-guez I, Nogueira JMF. 2012a. Combining stir-bar sorptive extraction and large volume injection-gas chromatography-mass spectrometry for the determination of benzotriazole UV stabilizers in wastewater matrices. J Sep Sci 35(3):459-467.

Carpinteiro I, AbuÃn B, Ramil M, RodrÃguez I, Cela R. 2012b. Matrix solid-phase dispersion followed by gas chromatography tandem mass spectrometry for the determination of benzotriazole UV absorbers in sediments. Analytical and Bioanalytical Chemistry 402(1):519-527.

Montesdeoca-Esponda S, Sosa-Ferrera Z, Santana-Rodríguez JJ. 2012 Mar. On-line solid-phase extraction coupled to ultra-performance liquid chromatography with tandem mass spectrometry detection for the determination of benzotriazole UV stabilizers in coastal marine and wastewater samples. Anal Bioanal Chem 2012(403):867-876.

Nakata H, Murata S, Filatreau J. 2009 Jula. Occurrence and Concentrations of Benzotriazole UV Stabilizers in Marine Organisms and Sediments from the Ariake Sea, Japan. Environ Sci Technol 43(18):6920-6926.

Nakata H, Sayaka M, Ryuichi S, Filatreau J, Isobe T, Takahashi S, Tanabe S. 2009 Marb. Occurrence and Concentrations of Persistent Personal Care Products, Organic UV Filters, in the Marine Environment. Interdisciplinary Studies on Environmental Chemistry 2:239-246.

Nakata H, Shinohara R. 2010 Jun. Concentrations of Benzotriazole UV Stabilizers and Polycyclic Musks in Wastewater Treatment Plant Samples in Japan. Int Stu Env Chem: 51-59.

Nakata H, Shinohara R, Murata S, Watanabe M. 2010 Aug. Detection of benzotriazole UV stabilizers in the blubber of marine mammals by gas chromatography-high resolution mass spectrometry (GC-HRMS). J Environ Monit(12):2088-2092.

Kameda Y, Kimura K, Miyazaki M. 2011. Occurrence and profiles of organic sunblocking agents in surface waters and sediments in Japanese rivers and lakes. Environmental Pollution 159(6):1570-1576.

Nakata H. 2011. Presentation: Benzotriazole UV Stabilizer (BUVS) in Human and Wildlife - Is it a POPs? 4th Inernational Conference on Environmental Health Science

- 2011, 27-28 October 2011 Seoul, Korea.

Nakata H, Shinohara RI, Nakazawa Y, Isobe T, Sudaryanto A, Subramanian A, Tanabe S, Zakaria MP, Zheng GJ, Lam PKS, Kim EY, Min BY, We SU, Viet PH, Tana TS, Prudente M, Frank D, Lauenstein G, Kannan K. 2012 Oct. Asia-Pacific mussel watch for emerging pollutants: Distribution of synthetic musks and benzotriazole UV stabilizers in Asian and US coastal waters. Marine Pollution Bulletin 64(10):2211-2218.

Yanagimoto H, Nakata H, Shinohara R, Isobe T, Tanabe S, Nose M, Komori H, Arita N, Ueda N, Watanabe M, Jemenez B, Yang J-H, Kunisue T, Kannan K. 2011. Poster: Occurrence of benzotriazole UV stabilizers and synthetic musks in human adipose tissues collected from Japan, South Korea, China, India, Spain, Poland and the USA. 32nd SETAC (Society of Environmental Toxicology and Chemistry) North America, Boston, USA, November 2011.

Nakata H, Murata S, Shinohara H, Yanagimoto H, Shikata N, Watanabe M, Isobe T, Tanabe S, Kannan K. 2011. Poster: Benzotriazole UV Stabilizers in the Environment: Is it a POPs? 32nd SETAC (Society of Environmental Toxicology and Chemistry) North America, Boston, USA, November 2011.

Watanabe M, Noma Y. 2010 Jun. Behavior of 2-(3,5-di-tert-butyl-2-hydroxyphenyl)benzotriazole (DBHPBT) and 2-(3,5-di-tert-butyl-2-hydroxyphenyl)-5-chlorobenzotriazole during incineration of solid waste contaminated with thousand mg/kg levels of DBHPBT. J Hazard Mater 178(1â€"3):1065-1069.

Kim JW, Ramaswamy BR, Chang KH, Isobe T, Tanabe S. 2011a. Multiresidue analytical method for the determination of antimicrobials, preservatives, benzotriazole UV stabilizers, flame retardants and plasticizers in fish using ultra high performance liquid chromatography coupled with tandem mass spectrometry. Journal of Chromatography A 1218(22):3511-3520.

Kim JW, Isobe T, Ramaswamy BR, Chang K-H, Amano A, Miller TM, Siringan FP, Tanabe S. 2011 Julb. Contamination and bioaccumulation of benzotriazole ultraviolet stabilizers in fish from Manila Bay, the Philippines using an ultra-fast liquid chromatography-tandem mass spectrometry. Chemosphere 85:751-758.

Kim JW, Isobe T, Malarvannan G, Sudaryanto A, Chang K-H, Prudente M, Tanabe S. 2012 Feb. Contamination of benzotriazole ultraviolet stabilizers in house dust from the Philippines: Implications on human exposure. Sci Total Environ:1-8.

Zhang Z, Ren N, Li YF, Kunisue T, Gao D, Kannan K. 2011 Apr. Determination of Benzotriazole and Benzophenone UV Filters in Sediment and Sewage Sludge. Environ Sci Technol 45:3909-3916.

Liu YS, Ying GG, Shareef A, Kookana RS. 2011b. Simultaneous determination of benzotriazoles and ultraviolet filters in ground water, effluent and biosolid samples using gas chromatography-tandem mass spectrometry. Journal of Chromatography A 1218(31):5328-5335.

Liu YS, Ying GG, Shareef A, Kookana RS. 2012. Occurrence and removal of benzotriazoles and ultraviolet filters in a municipal wastewater treatment plant. Environmental Pollution 165:225-232.

Jungclaus GA, Lopez-Avila V, Hites RA. 1978. Organic compounds in an industrial wastewater: A case study of their environmental impact. Environ Sci Technol 12(1):88-96.

Lopez-Avila V, Hites R. 1980. Organic compounds in an industrial wastewater. Their transport into sediments. Environ Sci Technol 14(11):1382-1390.

Pruell RJ, Hoffman EJ, Quinn JG. 1984. Total hydrocarbons, polycyclic aromatic hydrocarbons and synthetic organic compounds in the Hard shell clam, Mercenaria mercenaria, purchased at commercial seafood stores. Marine Environmental Research 11(3):163-181.

Reddy CM, Quinn JG, King JW. 2000. Free and Bound Benzotriazoles in Marine and Freshwater Sediments. Environ Sci Technol 34(6):973-979.

ANNEX 1: Read-Across-Data-Matrix

In this matrix all available experimental results that might be relevant for the SVHC-identification are listed for all four substances in questions as well as all other substances mentioned in the dossier or used for a Read Across. The substances are ordered in order of rising molecular weight.

QSAR results were intentionally left out in this overview. In cases where several data points were available the most reliable one is presented and in cases where a decisions was not possible (as is for example the case for registration data disseminated on ECHAs webpage) all data point are presented.

Acron ym	1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
		HO	CH C	но	B C C C C C C C C C C C C C C C C C C C	5						B B	
CAS No	95-14-7	2440- 22-4	3896- 11-5	3846- 71-7	3147- 75-9	36437- 37-3	84268- 36-0	25973- 55-1	3864- 99-1	125304 -04-3	73936- 91-1	70321- 86-7	103597 -45-1
EC No	202- 394-1	219- 470-5	223- 445-4	223- 346-6	221- 573-5	253- 037-1	-	247- 384-8	223- 383-8	-	422- 600-5	274- 570-6	403- 800-1
Physico	chemical D	ata		•	•	•		•	•		•	•	
Mol. Weigh t [g/mo I]	119.1	225.3	315.8	323.4	323.4	323.4	339.4	351.5	357.9	393.6	441.6	447.6	658.9
log	1.445	4.315										>6.56	4.27;

⁴ Degradation Product of EC 407-000-3

 ⁵ Hansch, C. et al: Exploring QSAR Vol 2: Hydrophobic, Electronic, and Steric Constants (1995)
 ⁶ The Phenolic Benzotriazoles Association: HPV Challenge Program, Data Summary and Test Plan for Phenoluic Benzotriazoles (2001)

1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
I Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	HO	A A A A A A A A A A A A A A A A A A A	HO HO HO HO HO HO HO HO HO HO HO HO HO H	B C C C C C C C C C C C C C C C C C C C	5							
	4.26											12.77,8
8.379												
												5.637
19800 10	0.173 11; 0.812			<16			0.0151 2	0.022 12			<0.04 (at 20°)6	<0.007 7
												6 10-13 7
n Degradati	ion											
non- biodegr adable MITI-1 (OECD TG	Not readily biodegr adable (OECD TG 301		non- biodegr adable MITI-1 (OECD TG	Not readily biodegr adable (OECD TG 301			Not readily biodegr adable (OECD TG 301	non- biodegr adable MITI-1 (OECD TG		Not readily biodegr adable (OECD TG 301	Not readily biodegr adable (OECD TG 301	Biodegr adation in water <10% (84/499 /CEE method
	Benzotr iazole	Benzotr iazole	Benzotr iazole Image: Constraint of the second state of the	Benzotr iazole Image: Construction of the second secon	Benzotr iazole Image: Construction of the second secon	Benzotr iazoleImage: second	Benzotr iazoleImage: second	Benzotr iazole Image: Construction of the second secon	Benzotr iazole Image: Second Seco	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Benzotr iazole Image: Second Seco	Benzotr iazole Image: Second Seco

 ⁷ Data disseminated on ECHA-Homepage
 ⁸ This value is so large that is probably not reliable
 ⁹ Serjeant,EP & Dempsey,B: Ionisation constants of organic acids in aqueous solution, p. 159 (1979)
 ¹⁰ Davis, LN et al: Investigation of selected potential environmental contaminants: benzotriazoles, USEPA-560/2-77-001 (1977)
 ¹¹ US EPA Screening-LevelHazard Characterization Sponsored Chemicals Phenolic Benzotriazoles Category (2009)
 ¹² Lopez-Avila, V & Hites, RA: EnvSciTechnol 11, p. 1382-1390 (1980)

Acron ym	1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
	I Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	HO		но	B B	5	A A A A A A A A A A A A A A A A A A A		Ho Look			5	A A A A A A A A A A A A A A A A A A A
	BOD =213	0-2% after 28 days11		BOD =013	0-1% after 28 days11			2-8% after 28 days11	BOD =013		-4-3% after 28 days14	3-8% after 28 days11	5) 7; Biodegr adation in water <2% (84/499 /CEE method 5) 7; Biodegr adation in water 0% (84/499 /CEE method 5) 7
Simul ation tests	Primary degrad ation aerobic : DT50= 114 d anaerob						OECD 308 aerobic : DisT50 = 86 d (river system)						

¹³ Biodegradation and Bioconcentration Database of the Existing Chemical Substances; available: <u>http://www.safe.nite.go.jp/jcheck/english/search.action</u> ¹⁴ Australia: Nantional Industrial Chemicals Notification and Assesment Scheme - Full Public Report - Tinuvin 928 (2000)

Acron ym	1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
	R N N N N N N N N N N N N N N N N N N N	HO	CH CH	HO HO	B C	B B C C C C C C C C C C C C C C C C C C			Ho Lo				Carter A
	ic: DT50= 144 d						DisT11 = 44 d (pond system) anaero bic: build up until test was ended (100 d)						
Data or BCF (lipid norma lized) acc. To OECD 305 C on Cyprin us carpio	Bioaccum 1000 μg/L: 1-3; 100 μg/L: 5-17 ¹³	ulation 1000 µg/L: 171- 686; 100 µg/L: 181- 410; 10 µg/L: 55- 275 ¹³	500 μg/L: 71-143; 50 μg/L: 258- 1055; 5 μg/L: 721- 1178 ¹³	10 µg/L: 1945; 1µg/L: 5905; 0.1µg/ L: 12041 ¹ ³		1 µg/L: 20263; 0.1 µg/L: 34210 ¹ ³		0.1 µg/L: 1121; 0.01 µg/L: 740- 2148; 0.01 µg/L: 3681 ¹³ 0,8 µg/L: 2655; 0,08 µg/L:	1 µg/L: 1203; 0.1µg/ L: 6283/ 8817; 0.01 µg/L: 7540 ¹³				

Acron ym	1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
	H N N N N N N N N N N N N N N N N N N N	HO	C C C C C C C C C C C C C C C C C C C	но	B C	B B C C C C C C C C C C C C C C C C C C	A A A A A A A A A A A A A A A A A A A		Contraction of the second seco				A A A A A A A A A A A A A A A A A A A
								5464 ¹⁵					
Field BAF calcul ated based on Nakat a et al 2010 on Neoph ocaen a phoca enoid									0.012 μg/L: 5946 ¹⁶				

¹⁵ Registration dossier of the lead registrant.
 ¹⁶: Nakata H et al.: Detection of benzotriazole UV stabilizers in the blubber of marine mammals by gas chromatography-high resolution mass spectrometry (GC-HRMS). J Environ Monit 12, p. 2088-2092 (2010)

Acron ym	1H- Benzotr iazole	UV-P	UV-326	UV-320	UV-329	UV-350	M14	UV-328	UV-327	UV-571	UV-928	UV-234	UV-360
		Ho Ho	A at a contract of the second	Ho the second se					A A A A A A A A A A A A A A A A A A A				A ANA
es													

ANNEX 2: Overview of Self-Classifications

Name / Tradename	EC-number	Hazard Class and Category Code(s)	Hazard Statement Code(s)
2-(2H-benzotriazol-2-yl)-	253-037-1	Eye Irrit. 2	H319
4-(tert-butyl)-6-(sec-		STOT RE 2	H373
butyl)phenol		Aquatic Chronic 4	H413

Table 17: Self Classification for UV-350 acc. to Regulation (EC) 1272/2008 (CLP)

ANNEX 3: Analysis of QSAR Application: Prediction of Potential persistent properties in the environment for UV-350

A Information on substance and purpose

Name:	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec- butyl)phenol (UV-350)	ОН
CAS Nr.	36437-37-3	
EU Nr.	253-037-1	\sim
Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)CC)O)N(N=C2C=C3) N=C2C=C3	*

Endpoint		Screening on Persistence in the environment
Regulatory	purpose	PBT-Assessment, used in a Weight-of-Evidence Approach to justify vP-
		assessment of a group of Phenolic Benzotriazoles

B Relevant structure information

D Relevant Structure								
Parameter	Result	Rationale						
Structure identification								
Structure of concern	parent	Substance is a mono-constituent, question to be answered is, if this substance itself fulfills the P-Screening criteria.						
Descriptors used for QS	SAR prediction							
Fragment descriptors of BIOWIN2 and BIOWIN 3		All fragments are represented by the model with exception of the triazole group which has probably an inhibiting influence						
Fragment descriptors of BIOWIN6	applicable but probably slightly underestimating	All fragments are represented by the model with exception of the triazole group which has probably an inhibiting influence.						
Molecular Weight	applicable	Molecular weight in the range of the weights in the Training Set						
Other relevant information								
-	-	-						

C QSAR models used

Model	Version	Endpoint	QMBI
BIOWIN2	v4.1	Probability of rapid biodegradation	Annex 3.1
BIOWIN3	v4.1	Time to Ultimate Biodegradation given as a numerical value between 1 and 5	Annex 3.2
BIOWIN6	v4.1	Probability of rapid biodegradation	Annex 3.3

D Analysis of QSAR model performance

Model	QSAR result	Overall model performance	QPREF
BIOWIN2	0.1329 (does not biodegrade fast)	Reliable with Restrictions	Annex 3.4
BIOWIN6	0.012 (does not biodegrade fast)	Reliable with Restrictions	Annex 3.4

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BIOWIN3	2.2538 (weeks	Reliable with Restrictions	Annex 3.4
	to months	5)		

E Overall conclusion

Overall QSAR Result	Borderline case for meeting the criteria on persistence
Rational	Both QSAR-criteria according to ECHA Guidance R.11 are not met ((BIOWIN2 < 0.5 AND BIOWIN3 < 2.2) OR (BIOWIN6 < 0.5 AND BIOWIN 3 < 2.2))). The reason for this is that the BIOWIN3-result is larger by 0.0538 which indicates a borderline case The model performance and Read-Across on similar substances supports the result of the QSAR model (see Annex 3.4).
Reliability	Reliable with Restrictions. The triazole group is not a descriptor fragment of the three models, therefore the actual persistence might be underestimated. The comparison of experimental data for similar structures with BIOWIN3 results for them can be interpreted as showing a slight underestimation of the Ultimate Biodegradation time, therefore the actual degradation time might probably be longer and the criterion fulfilled.

Conclusion with regard to the regulatory purpose When combining the evidence of the QSAR results with the information known from similar structures it can be concluded that UV-350 meets the Persistence criterion on a Screening level. Note that the information obtained in this way is on its own not sufficient for assessing the actual P/vP-criterion. This will be done in a Weight-of-Evidence approach where this QSAR result is used as supporting evidence.

ANNEX 3.1: QMBI BIOWIN2

	Information	Literature references or Links	Remarks			
0 - General	J - General					
Model name and version	BIOWIN2 (nonlinear probability model), v4.10	 Howard, P.H., Boethling, R.S., Stiteler, W.M., Meylan, W.M., Hueber, A.E., Beauman, J.A., Larosche, M.E. 1992. Predictive model for aerobic biodegradability developed from a file of evaluated biodegradation data. Environ. Toxicol. Chem. 11:593-603. Boethling, R.S., Howard, P.H., Meylan, W., Stiteler, W., Beaumann, J., Tirado, N. 1994. Group contribution method for predicting probability and rate of aerobic biodegradation. Environ. Sci. Technol. 28:459-65. 	-			
w.a.17: software package	EPISUITE Estimation Programs Interface Suite [™] for Microsoft® Windows, v4.10	http://www.epa.gov/oppt/exposure/pubs/episuite.htm	-			
1 - Definition of Endp	oint					
Endpoint [units] (w.a. species and other relevant information)	Probability of Rapid Degradation (0.0-1.0)		Values larger 0.5 are interpreted as biodegradades fast			
2 – Definition of Algor	ithm					
Brief description of algorithm and/or link to full definition		Online documentation of BIOWIN	-			
List of employed descriptors with units	a0=3.0087; a1a36: fragment coefficients, f1f36: instances of fragment, am: molecular weight coefficient, MWj : molecular weight of substance j [g/mol]	List of the 36 fragments employed and all their coefficients can be found in online documentation of BIOWIN	-			
Number of Chemicals in	295 chemicals of the BIODEG database		-			

¹⁷w.a.: when applicable

Training Set and Brief description of it			
W.a.: Training set available at		Online documentation of BIOWIN	
3 – Definition of the A	pplicability Domain		
W.a.: Definition of the Applicability Domain	-	-	-
Limits of the Applicability Domain	The target molecule should be composed of fragments that are used in this model, otherwise only molecular weight will be considered; MW = 30.02-959.2	Online documentation of BIOWIN	-
	e Validation of the Model		
Validation Set Type	internal and external		-
W.a.: Validation available at		Several available, see for example: Posthumus, R., Traas, T.P., Peijnenburg, W.J.G.M., Hulzebos, E.M. 2005. External validation of EPIWIN biodegradation models. SARQSAR Environ. Res. 16:135-148	-
Statistical information on validity	Total correct: 275 / 295 (93.2%)		-

5 – Mechanistic Inter	5 – Mechanistic Interpretation of the model			
W.a.: Mechanistic	Biodegradation is a	-	-	
basis of model	stepwise reaction			
	cascade therefore			
	individual fragments			
	contribute to the			
	probability that a			
	substance is degraded			
	fast			

ANNEX 3.2: QMBI BIOWIN3

	Information	Literature references or Links	Remarks
0 - General			
Model name and version	BIOWIN3 (ultimate survey model), v4.10	Boethling, R.S., Howard, P.H., Meylan, W., Stiteler, W., Beaumann, J., Tirado, N. 1994. Group contribution method for predicting probability and rate of aerobic biodegradation. Environ. Sci. Technol. 28:459-65.	-
w.a.18: software package	EPISUITEEstimationProgramsInterfaceSuite™forMicrosoft®Windows, v4.10	http://www.epa.gov/oppt/exposure/pubs/episuite.htm	
1 - Definition of Endp	oint		
Endpoint [units] (w.a. species and other relevant information)	Expert Rating for time to Ultimate Biodegradation given as a numerical value between 1 and 5		>4.75 - 5: hours; >4.25 - 4.75: hours to days; >3.75 - 4.25: days; >3.25 - 3.75: days to weeks; >2.75 - 3.25: weeks; >2.25 - 2.75: weeks to months; >1.75 - 2.25: months; <1.75: recalcitrant
2 – Definition of Algor			
Brief description of algorithm and/or link to full definition	a2 * f2 + + a36 * f36 + am * MWj	Online documentation of BIOWIN	-
List of employed descriptors with units	a0=3.1992; a1a36: fragment coefficients, f1f36: instances of fragment, am: molecular weight coefficient, MWj : molecular weight of	List of the 36 fragments employed and all their coefficients can be found in online documentation of BIOWIN	-

¹⁸w.a.: when applicable

	substance j [g/mol]		
Number of Chemicals in Training Set and Brief description of it	200 substances		17 Experts reviewed 200 substances and assigned a degradation rating of 1 to 5 for them. Based on this rating the coefficients for the fragments
			were derived
W.a.: Training set available at		Online documentation of BIOWIN	-
3 – Definition of the A	policability Domain		
W.a.: Definition of	-	-	-
the Applicability Domain			
Limits of the Applicability Domain	The target molecule should be composed of fragments that are used in this model, otherwise only molecular weight will be considered; Prediction Range for Expert Rating : 1.44 – 3.89	Online documentation of BIOWIN	-
	e Validation of the Model		
Validation Set Type	internal and external		-
W.a.: Validation available at		Several available, see for example: Posthumus, R., Traas, T.P., Peijnenburg, W.J.G.M., Hulzebos, E.M. 2005. External validation of EPIWIN biodegradation models. SARQSAR Environ. Res . 16:135-148	-
Statistical information on validity	Total correct: 167/200 (83.5%)		-
5 – Mechanistic Interp			
W.a.: Mechanistic	Biodegradation is a	-	-

basis of model	stepwise reaction cascade therefore
	individual fragments contribute to the
	probability that a substance is degraded
	fast

ANNEX 3.3: QMBI BIOWIN6

	Information	Literature references or Links	Remarks		
0 - General	0 - General				
Model name and version	BIOWIN6 (non-linear MITI model), v4.10	Tunkel, J., Howard, P.H., Boethling, R.S., Stiteler, W., Loonen, H. 2000. Predicting Ready Biodegradability in the Japanese Ministry of International Trade and Industry Test. Environ. Toxicol. Chem. 19:2478-2485.	-		
w.a.19: software package	EPISUITE Estimation Programs Interface Suite [™] for Microsoft® Windows, v4.10	http://www.epa.gov/oppt/exposure/pubs/episuite.htm	-		
1 - Definition of Endp	point				
Endpoint [units] (w.a. species and other relevant information)	Probability of Rapid Degradation (0.0-1.0)		Values larger 0.5 are interpreted as "biodegradades fast"		

2 – Definition of Algor	2 – Definition of Algorithm			
Brief description of	Probability = Exp (Yj) /	Online documentation of BIOWIN	-	
algorithm and/or	(1 + Exp (Yj)), Yj = a0 +			
link to full definition	a1* f1 + a2 * f2 + +			
	a42 * f42 + am * MWj			
List of employed	a0= 2.5257; a1a42:	List of the 42 fragments employed and all their coefficients can be	-	
descriptors with	fragment coefficients,	found in online documentation of BIOWIN		
units	f1f42: instances of			
	fragment, am: molecular			
	weight coefficient, MWj :			
	molecular weight of			
	substance j [g/mol]			
Number of	589 chemicals of 884		The other 295	
Chemicals in	chemicals that were		substances were	
Training Set and	tested under the		used as Validation	
Brief description of	Japanese Chemical		Set	
it	Substances Control Law			
W.a.: Training set		Online documentation of BIOWIN	-	

¹⁹w.a.: when applicable

available at				
3 – Definition of the Applicability Domain				
W.a.: Definition of	-	-	-	
the Applicability				
Domain				
Limits of the	The target molecule	Online documentation of BIOWIN	-	
Applicability Domain	should be composed of			
	fragments that are used			
	in this model, otherwise			
	only molecular weight			
	will be considered			
	e Validation of the Model			
Validation Set Type	internal and external		-	
W.a.: Validation		Internal: Online documentation of BIOWIN	-	
available at		External: Several available, see for example:		
		Posthumus, R., Traas, T.P., Peijnenburg, W.J.G.M., Hulzebos, E.M.		
		2005. External validation of EPIWIN biodegradation models. SARQSAR		
		Environ. R e s . 1 6 : 1 3 5 - 1 4 8		
Statistical	Total correct Training		-	
information on	Set: 488 / 589 (82.9%)			
validity	Total Correct Validation			
	Set: 238 / 295 (80.7%)			
5 – Mechanistic Interp				
W.a.: Mechanistic	Biodegradation is a		-	
basis of model	stepwise reaction			
	cascade therefore			
	individual fragments			
	contribute to the			
	probability that a			
	substance is degraded			
	fast			

ANNEX 3.4: Analysis of QSAR prediction for UV-350

QSAR Model: BIOWIN 2/3/6

Overall performance

	Result	Further description
Endpoint result [unit]	BIOWIN2: 0.1329 BIOWIN6: 0.012 BIOWIN3: 2.2538 Borderline Case of Persistence Criterion met	According to the ECHA Guideline R.11 the Screening Criterion is met, when (BIOWIN2 < 0.5 AND BIOWIN3 < 2.2) OR (BIOWIN6 < 0.5 AND BIOWIN3 < 2.2), this is not the case here since the BIOWIN3 result is 0.0538 above the trigger value.
Applicability domain	Yes, with restrictions	The molecule is in the range of molecular weights of the training sets of the three models. Also, most of the fragments of UV-350 are represented in the fragment descriptors of the molecule (with the exception of the triazole group). The prediction of BIOWIN 3 is in the range of the predictions of the Training Set as well.
Similarity with trainings set	Yes	Most fragments of UV-350 are present in the Training Set (with the exception of the triazole group)
Similar substances	Yes	See table next side
Model performance for similar substances	Good	Experimental Values and Assessment are in agreement
Other uncertainties	No	-

Overall conclusion	Reliable with restrictions (but used in a Weight of Evidence Approach)
Rational	The result is very near the trigger value for BIOWIN3 (< 2.2). One essential group for the degradation, the triazole group is not represented. As the group is known to be difficult to degrade its contribution should probably be inhibiting. Hence, it can be assumed that UV-350 will be indeed not readily-biodegradable. This assumption is supported by the known experimental values of other phenolic benzotriazoles. The comparison of experimental data for similar structures with BIOWIN3 results for them can be interpreted as showing a slight underestimation of the Ultimate Biodegradation time.

Results for similar substances

	Substance 1	Substance 2	Substance 3	Substance 4	Substance 5
Structure		OH N N	OH N N		C) C) C) C) C) C) C) C) C) C) C) C) C) C
CAS-Nr.	3864-99-1	3846-71-7	25973-55-1	3147-75-9	70321-86-7
EU-Nr.	223-383-8	223-346-6	247-384-8	221-573-5	274-570-6
(Trade-)Name	UV-327	UV-320	UV-328	UV-329	UV-234
Descriptor value	BIOWIN 2 : 0.0013; BIOWIN6 : 0.0024; BIOWIN 3: 1.8338 (months)	BIOWIN 2: 0.016; BIOWIN6: 0.0091; BIOWIN 3: 2.1165 (months)	BIOWIN 2 : 0.0108; BIOWIN6 : 0.0096; BIOWIN 3: 2.0546 (months)	BIOWIN 2 : 0.016; BIOWIN6 : 0.0154; BIOWIN 3: 2.1165 (months)	BIOWIN 2 : 0.0922; BIOWIN6 : 0.0008 ; BIOWIN 3: 1.8862 (months)
Predicted endpoint	P-Screening criterion met	P-Screening criterion met	P-Screening criterion met	P-Screening criterion met	P-Screening criterion met
Experimental endpoint	Non-biodegradable, BOD = 0	Non-biodegradable, BOD = 0	not readily biodegradable (2- 8% after 28 days)	not readily biodegradable (0- 1% after 28 days)	not readily biodegradable (3- 8% after 28 days)
Statistical performance	-	-	-	-	-

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	Substance 6
Structure	
CAS-Nr.	73936-91-1
EU-Nr.	403-800-1
(Trade-)Name	UV-928
Descriptor value	BIOWIN 2 : 0.0027; BIOWIN6 : 0.0013 ; BIOWIN 3: 1.6343 (recalcitrant)
Predicted endpoint	P-Screening criterion met
Experimental endpoint	not readily- biodegradable (0- 3% after 28 days)
Statistical performance	-

Rational for the selection of similar substances

Substances 1 to 6 are structurally similar as they are phenolic benzotriazoles as the target molecule. The largest similarity is with Substance 2 and 3.

ANNEX 4: Analysis of QSAR Application: Prediction of log KOC for UV-320, -327, -328 and -350

A Information on substances and purpose

Molecule 1:

Name:	2-benzotriazol-2-yl-4,6-di-tert-butylphenol (UV- 320)	ОН		
CAS Nr.	3846-71-7			
EU Nr.	223-346-6	\mathbf{X}		
Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)(C)C)O)N(N=C2C=C 3)N=C2C=C3			

Molecule 2:

Name:	2,4-di-tert-butyl-6-(5-chlorobenzotriazol-2- yl)phenol (UV-327)	ОН
CAS Nr.	3864-99-1	
EU Nr.	223-383-8	
Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)(C)C)O)N(N=C2C=C 3)N=C2C=C3Cl	

Molecule 3:

Name:	2-(2H-benzotriazol-2-yl)-4,6-ditertpentylphenol (UV-328)	он
CAS Nr.	25973-55-1	
EU Nr.	247-384-8	\times
Smiles	c1(c(c(cc(c1)C(C)(C)CC)C(C)(C)CC)O)N(N=C2C =C3)N=C2C=C3	,

Molecule 4:

Name:	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec- butyl)phenol (UV-350)	ОН
CAS Nr.	36437-37-3	
EU Nr.	253-037-1	

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Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)CC)O)N(N=C2C=C3) N=C2C=C3			
Endpoint	pint Logarithmic Partition coefficient of octanol-organic carbon			
Regulatory	purpose	PBT-Assessment, supporting information for a weight of evidence-		
approach to identify the substances as vP				

B Relevant structure information

Parameter	Result	Rationale		
Structure identification				
Structure of concern	parent	Substances are mono-constituents		
Descriptors used for QS	SAR prediction			
Correction factors (KOCWIN KOW/MCI)	Applicable	All fragments are represented by the model		
□ (COSMOtherm)	Applicable	The polarity was calculated on molecular structures geometrically optimized with Density-Functional-Theory (functional: Becke- Perdew 86, basis set of Triple-Zeta-Valence- Polarization-quality), all parameters for this method and all elements of the molecules are implemented		
Other relevant information				
-	-	-		

C QSAR models used

Model		Version	Endpoint	QMBI
(PC)KOCWIN	-	V2.0	log KOC	Annex 4.1
KOW method				
(PC)KOCWIN	-	V2.0	log KOC	Annex 4.2
MCI method				
COSMOtherm		v.	log KOC	Annex 4.3
(KOC)		C30_1201		

D Analysis of QSAR model performance

Model	QSAR result	Overall model performance	QPREF
KOCWIN KOW	UV-320: 4.63	Reliable with restrictions	Annex 4.4
method	UV-327: 4.99		
	UV-328: 5.18		
	UV-350: 4.66		
KOCWIN MCI	UV-320: 5.07	Reliable with restrictions	Annex 4.4
method			
	UV-327: 5.28		
	UV-328: 5.65		
	UV-350: 5.19		
COSMOtherm	UV-320: 5.17	Reliable with restrictions	Annex 4.4
(KOC)	UV-327: 5.64		
	UV-328: 5.46]	
	UV-350: 4.90		

E Overall conclusion

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Overall QSAR Result	Irrespective of the employed model all four substances have a high log KOC. There does not seem to be a general systematic shift between the models and there is also no general order of the values when comparing the relative order of the results in the three models.
Rational	The log KOC for all substances and all models is in the range of 4.63 to 5.65 log-units
Reliability	Reliable with restrictions.

Conclusion with regard to the regulatory purpose The log KOC-values for all four substances are high in all three models. The predictions are all in the same region, therefore these substances are similar in their behavior. According to the prediction the substances will bind strongly to sediment in the environment and therefore will mostly not be available for degradation processes.

	Information	Literature references or Links	Remarks
0 - General			
Model name and version	(PC)KOCWIN v.2 - KOW method	Online Help of KOCWIN	The KOCWIN – KOW method is essentially an extension of the MCI method were the descriptor MCI was replaced with KOW. The same Trainings Sets and Validation Sets as for the MCI method were used and also the same Correction factors are applied. Overall the statistical performance of the KOW method is not quite as good as the MCI method.
W.a.20: software package	EPISUITE Estimation Programs Interface Suite [™] for Microsoft® Windows, v4.10	http://www.epa.gov/oppt/exposure/pubs/episuite.htm	
1 - Definition of			
Endpoint [units] (w.a. species and other relevant information)	Soil adsorption coefficient KOC given as a logarithmic value		Definition of KOC according to Lyman et al, 1990: "the ratio of the amount of chemical adsorbed per unit weight of organic carbon (oc) in the soil or sediment to the concentration of the chemical in solution at equilibrium" Koc = (µg adsorbed/g organic carbon) / (µg/mL solution) [L/kg or mL/g]
2 – Definition of			The
Brief	Non-polar chemicals (i.e.	See Online Help of KOCWIN	The equations were

²⁰w.a.: when applicable

description of algorithm and/or link to full definition	compounds where no correction factor is needed): log Koc = 0.8679 Log Kow - 0.0004 Polar chemicals (i.e. compounds where a correction factor is needed): log Koc = 0.55313 Log Kow + $0.9251 + \Sigma PfN$		developed in a two separate regression calculations since this approach is statistically more accurate than the approach taken in the MCI- method
List of employed descriptors with units	Log KOW: logarithm of the n- octanol/water partition coefficient; Pf: correction factor for chemical class of functional group f; N: number of times chemical class or functional group f occurs	List of Pf available in Online Help of KOCWIN, Appendix D	
Number of Chemicals in Training Set and Brief description of it	Training Set comprises of non-polar set (68 chemicals) and a polar set (447 chemicals) taken from several literature sources. One compound of the original non-polar training set (hexabromobiphenyl) was not considered since there was no recommended experimental log KOW.		Training Estimation Error: within <= $0.20 - 44.2\%$ within <= $0.40 - 76.9\%$ within <= $0.60 - 93.0\%$ within <= $0.80 - 98.6\%$ within <= $1.00 - 100\%$ non-polar Training Set (n=68): r2= 0.877 ; std. dev.= 0.478 ; avg. dev.= 0.371 polar Training Set (n=447): r2= 0.855 ; std. dev.= 0.396 ; avg. dev.= 0.307
W.a.: Training set available at		Non-Polar Training Set: Online Help of KOCWIN, Appendix E Polar Set: Online Help of KOCWIN, Appendix F	
3 – Definition of	f the Applicability Domain		
W.a.: Definition of the Applicability Domain	Currently there is no universally	KOCWIN, Appendix D Non-Polar Training Set: Online Help of KOCWIN, Appendix E	

	compounds and/or that have more instances of a given fragment than the maximum for all training set compounds. It is also possible that a compound may have a functional group(s) or other structural features not represented in the training set, and for which no fragment coefficient or correction factor was		
Limits of the	developed		
Applicability	Molecular weight: 32.04-665.02 g/Mol		
Domain	Fragments and Functional groups		
	according to Training Sets and		
	correction factors for best results		
	on the Validation of the Model		
Validation Set	Internal, 150 compounds from the same sources as the Training Set.		
Туре	Eight ammonium and metal salt		
	compounds were removed from the		
	original Validation dataset of the		
	MCI method. Compound Pool was		
	split before regression into Training		
	Set and Validation Set.		
W.a.:		Online Help of KOCWIN, Appendix G	
Validation available at			
Statistical	r2=0.778; std. dev.=0.679; avg.		
information on	dev.= 0.494		
validity			
5 – Mechanistic Interpretation of the model			
W.a.:	The tendency of a compound to		
Mechanistic	adsorb itself on organic carbon is		
basis of model	linked with its lipophilicity. The n-		
	octanol/water partition coefficient is		
	one descriptor for lipophilicity.		

ANNEX 4.2: QMBI KOCWIN MCI-method

	Information	Literature references or Links	Remarks
0 – General			
Model name and version	(PC)KOCWIN v.2 - MCI method	Meylan, W., P.H. Howard and R.S. Boethling, "Molecular Topology/Fragment Contribution Method for Predicting Soil Sorption Coefficients", Environ. Sci. Technol. 26: 1560-7 (1992)	Besides the MCI method there is also the KOW method implemented in KOCWIN. Overall the statistical performance of the MCI method is better than the KOW method.
W.a.21: software	EPISUITE Estimation Programs Interface Suite [™] for Microsoft®	http://www.epa.gov/oppt/exposure/pubs/episuite.htm	
package 1 - Definition of	Windows, v4.10		
			Defintion of KOC
Endpoint [units] (w.a. species and other relevant information)	Soil adsorption coefficient KOC given as a logarithmic value		Definition of KOC according to Lyman et al, 1990: "the ratio of the amount of chemical adsorbed per unit weight of organic carbon (oc) in the soil or sediment to the concentration of the chemical in solution at equilibrium" Koc = (µg adsorbed/g organic carbon) / (µg/mL solution) [L/kg or mL/g]
2 – Definition of			
Brief description of algorithm and/or link to full definition	log Koc = 0.5213 MCI + 0.60 + Σ(Pf*N); MCI = Σ(δi*δj)-0.5	See Online Help of KOCWIN	MCI: Molecular Connectivity Index (in this case: First Order) mathematical approach to describe molecular topology The equation was

²¹w.a.: when applicable

			 developed in a two step regression approach: 1. Derivation of equation without correction factors using a set of non polar chemicals 2. Derivation of final equation using a set of non-polar chemicals
List of employed descriptors with units	δ i: δ-value of atom i, i.e. the number of adjacent non- hydrogen atoms; δ j: δ value of atom j, i.e. the number of adjacent non-hydrogen atoms; Pf: correction factor for chemical class of functional group f; N: number of times chemical class or functional group f occurs	List of Pf available in Online Help of KOCWIN, Appendix D	
Number of Chemicals in Training Set and Brief description of it	Training Set comprises of non- polar set (69 chemicals) and a polar set (447 chemicals) taken from several literature sources		Training Set Estimation Error: within <= 0.20 - 44.2% within <= 0.40 - 76.9% within <= 0.60 - 93.0% within <= 0.80 - 98.6% within <= 1.00 - 100% non-polar Training Set (n=69): r2=0.967; std. dev.=0.247; avg. dev.= 0.199 polar Training Set (n=447): r2=0.90; std. dev.=0.34; avg. dev.=

W.a.: Training		Non-Polar Training Set: Online Help of KOCWIN, Appendix E			
set available at		Polar Set: Online Help of KOCWIN, Appendix F			
	3 – Definition of the Applicability Domain				
W.a.: Definition of the Applicability Domain	Currently there is no universally accepted definition of model domain. Log Koc estimates are less accurate for compounds outside the MW range of the training set compounds and/or that have more instances of a given fragment than the maximum for all training set compounds. It is also possible that a compound may have a functional group(s) or other structural features not represented in the training set, and for which no fragment coefficient or correction factor was developed	List of correction factors available in Online Help of KOCWIN, Appendix D Non-Polar Training Set: Online Help of KOCWIN, Appendix E Polar Training Set: Online Help of KOCWIN, Appendix F			
Limits of the Applicability Domain	Molecular weight: 32.04-665.02 g/Mol Fragments and Functional groups according to Training Sets and correction factors for best results				
4 – Information	on the Validation of the Model				
Validation Set Type					
W.a.: Validation available at		Online Help of KOCWIN, Appendix G			
Statistical information on validity	_				
5 – Mechanistic	Interpretation of the model				

W.a.: Mechanistic basis of model	The tendency of a compound to adsorb itself on organic carbon is linked with the chemical structure. In the Molecular Correction Index information on the chemical structure, i.e. molecular size, branching, cyclization, unsaturation and (to a certain extent) heteroatom content are encoded. The different influences of chemical classes or functional groups are	
	or functional groups are considered by correction	
	factors.	

ANNEX 4.3: QMBI COSMOtherm (KOC)

	Information	Literature references or Links	Remarks
0 - General			
Model name and version	COSMOtherm v C30_1201		The COSMOtherm model allows in principle the calculation of all partition properties of molecules. In this QMBI only the calculation of the KOC will be addressed
W.a.22: software package	COSMOtherm		
1 - Definition of I	Endpoint		
Endpoint [units] (w.a. species and other relevant information)	n-octanol/organic carbon partition coefficient given as a logarithmic value		
2 – Definition of A	Algorithm		
Brief description of algorithm and/or link to full definition	Log KOC = $0.0168*M0X - 0.017*M2X - 0.040*M3X + 0.19*PMaccX - 0.27*MdonX + 0.37 with MiX = \int p^X \sigma^i d\sigma for i = 0, 2, 3. M_{acc}^X = 0 if \sigma < 1 \text{ e/nm}^2 or = \sigma - 1 \text{ e/nm}^2 if \sigma > 1 \text{ e/nm}^2 and M_{don}^X = 0 if -\sigma < 1 \text{ e/nm}^2 or = -\sigma - 1 \text{ e/nm}^2 if -\sigma > 1 \text{ e/nm}^2$	"COSMO-RS: From Quantum Chemistry to Fluid Phase Thermodynamics and Drug Design", Andreas Klamt, Elsevier Science Ltd., Amsterdam, The Netherlands (2005), ISBN: 0- 444-51994-7. "Prediction Of Soil Sorption Coefficients With A Conductor- Like Screening Model For Real Solvents", Andreas Klamt, Frank Eckert and Michael Diedenhofen, Environmental Toxicology and Chemistry, 21, 2562-2566 (2002).	COSMOtherm implements the COSMO-RS theory. This theory interprets the interaction of molecules as an interaction of a larger ensemble of molecular surfaces calculated with Quantum Mechanical methods. Due to a treatment with statistical thermodynamics the macroscopic properties of interacting molecules like partition coefficients become available. If the partition is with a phase that is ill defined like organic carbon, the so called σ -moment approach is employed where a solvent is represented as a linear combination of six σ -functions. The coefficients to these functions are fitted with experimental data.
List of	σ : Screening charge density or polarity,		

²²w.a.: when applicable

employed descriptors with units Number of	i.e. the electrostatic screening of a solute molecule by its surrounding and its back polarization in a region with radius of ca. 0.5 Å; pX: sigma profile of molecule X, i.e. the sum of the probability distributions of all possible σ Original parameterization for		While the principle theory is applicable for
Chemicals in Training Set and brief description of it	COSMOtherm: 225 small- and medium- sized organic compounds with H, C, O, N, Cl atoms. The fitting was done for 650 experimental room-temperature parameters (Δ Ghydr, log(vapor pressure), log Koctanol-water, log Khexane-water, log Kbenzene-water, log Kdiethyl ether-water log KOC-formula: 387 molecules (performance: r2 = 0.72, rms = 0.62 log-units)		all elements, the practical implementation needs some specific parameters to the QM- method used and the elements of the substance in question like the employed ratio for scaling the bonds of the QM- method and the van der Waals-coefficients
W.a.: Training set available at	ha Anglianhilitu Damain	Original parameterization for COSMOtherm: "Refinement and Parametrization of COSMO-RS", Andreas Klamt, Volker Jonas, Thorsten Bürger and John C. W. Lohrenz, J. Phys. Chem. A 102, 5074-5085 (1998). Log KOC-formula: "Prediction Of Soil Sorption Coefficients With A Conductor- Like Screening Model For Real Solvents", Andreas Klamt, Frank Eckert and Michael Diedenhofen, Environmental Toxicology and Chemistry, 21, 2562-2566 (2002).	Original parameterization for COSMOtherm: Since the original parameterization was done further adjustments were made and parameters for further elements were introduced. While the parameters are available in the software, to our knowledge the details of the new parameterisations were not disclosed
	he Applicability Domain There is no formal definition of the		
of the Applicability	applicability domain		

Domain			
Limits of the	In principle the method is completely		
Applicability	based on first-principles meaning there		
Domain	is no limit of the Applicability Domain.		
	n the Validation of the Model		
	The KOC-model was tested against 53 demanding chemicals achieving a rmd of 0.72		
W.a.: Validation available at		"Prediction Of Soil Sorption Coefficients With A Conductor- Like Screening Model For Real Solvents", Andreas Klamt, Frank Eckert and Michael Diedenhofen, Environmental Toxicology and Chemistry, 21, 2562-2566 (2002).	
Statistical	-		
information on			
validity			
	nterpretation of the model		
W.a.:	The interaction of a solute and a solvent		
Mechanistic	is calculated in terms of a chemical		
basis of model	potential. The difference of the chemical		
	potentials of the solute in two different		
	solvents is the mechanistic reason for partition effects.		

ANNEX 4.4: Analysis of QSAR prediction for UV-320 , UV-327, UV-328, UV-350

QSAR Model: KOCWIN KOW-method, KOCWIN MCI-method and COSMOtherm (KOC)

Overall performance

	Result		Further description
Endpoint results	KOCWIN	UV-320: 4.63	All log KOC-values are high and in a
[unit]	KOW-method	UV-327: 4.99	similar region.
		UV-328: 5.18	
		UV-350: 4.66	
	KOCWIN	UV-320: 5.07	
	MCI-method	UV-327: 5.28	
		UV-328: 5.65	
		UV-350: 5.19	
	COSMOther	UV-320: 5.17	
	m (KOC)	UV-327: 5.64	
		UV-328: 5.46	
		UV-350: 4.90	
Applicability domain	Yes		The molecules are in the range of all
			descriptors employed in the models.
Similarity with	Yes		All fragments or elements of the
trainings set			molecules are represented in the
			Training Set of KOCWIN.
			COSMOtherm has no training set but
			is generally applicable.
Similar substances	One		See table next side, substance is not
			very similar
Model performance	Mediocre		There is just one experimental value
for similar			of unknown quality for a substance
substances			not very similar to the substances at
			hand. The prediction for this
			substance is much higher than the
			experimental value but both values
			are high.
Other uncertainties	No		-

Overall conclusion	Reliable
Rational	As the models are applicable and results for similar molecules and two of the four models at hand show values in the same range it can be expected that the range is correctly predicted.

Results for similar substances

	Substance 1
Structure	
CAS-Nr.	103597-45-1
EU-Nr.	403-800-1

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(Trade-)Name	UV-360
Descriptor value	KOCWIN KOW-method : log KOC = 11.08 KOCWIN KOW-method : log KOC = 8.22 COSMOtherm: log KOW = 7.91
Predicted endpoint	See above
Experimental endpoint	5.63
Statistical performance	-

Rationale for the selection of similar substances

Substance 1 is a phenolic benzotriazole as the target molecule but it is a molecule comprised of two phenolic benzotriazole bodies therefore the similarity is not very high. Since the functional groups are nevertheless the same and since there are no other phenolic benzotriazoles were a experimental log KOC is reported, UV-360 was chosen as point of reference.

ANNEX 5: Analysis of QSAR Application: Prediction of log KOW for UV-320, -327, -328 and -350

A Information on substances and purpose

Molecule 1:

Name:	2-benzotriazol-2-yl-4,6-di-tert-butylphenol (UV- 320)	ОН		
CAS Nr.	3846-71-7			
EU Nr.	223-346-6	\mathbf{X} " $\mathbf{\tilde{x}}$		
Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)(C)C)O)N(N=C2C=C 3)N=C2C=C3			

Molecule 2:

Name:	2,4-di-tert-butyl-6-(5-chlorobenzotriazol-2- yl)phenol (UV-327)	ОН
CAS Nr.	3864-99-1	
EU Nr.	223-383-8	
Smiles	c1(c(c(cc(c1)C(C)(C)C)C(C)(C)C)O)N(N=C2C=C 3)N=C2C=C3Cl	

Molecule 3:

Name:	2-(2H-benzotriazol-2-yl)-4,6-ditertpentylphenol (UV-328)	он
CAS Nr.	25973-55-1	
EU Nr.	247-384-8	\times
Smiles	c1(c(c(cc(c1)C(C)(C)CC)C(C)(C)CC)O)N(N=C2C =C3)N=C2C=C3	,

Molecule 4:

Name:	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec- butyl)phenol (UV-350)	ОН
CAS Nr.	36437-37-3	
EU Nr.	253-037-1	

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Smiles	c1(c(c(cc(c N=C2C=C3	c1)C(C)(C)C)C(C)CC)O)N(N=C2C=C3) 3		
Endpoint		Logarithmic Partition coefficient of octanol-water		
Regulatory	purpose	purpose PBT-Assessment, supporting information		

B Relevant structure information

Parameter	Result	Rationale	
Structure identification			
Structure of concern	parent	Substances are mono-constituents	
Descriptors used for QS	SAR prediction		
Fragment descriptors (KOWWIN)	applicable	All fragments are represented by the model	
σ (COSMOtherm)	applicable	The polarity was calculated on molecular structures geometrically optimized with employing Density-Functional-Theory (functional: Becke-Perdew 86, basis set of Triple-Zeta-Valence-Polarization-quality), all parameters for this method and all elements of the molecules are implemented	
Other relevant information			
-	-	-	

C QSAR models used

Model	Version	Endpoint	QMBI
KOWWIN	v1.68	log KOW	Annex 5.1
COSMOtherm	v.	log KOW	Annex 5.2
(KOW)	C30_1201	_	

D Analysis of QSAR model performance

Model	QSAR result	Overall model performance	QPREF
KOWWIN	UV-320: 6.27	Reliable	Annex 5.3
	UV-327: 6.91		
	UV-328: 7.25		
	UV-350: 6.31		
COSMOtherm	UV-320: 7.39	Reliable	Annex 5.3
(KOW)	UV-327: 7.91		
	UV-328: 7.89		
	UV-350: 7.11		

E Overall conclusion

Overall QSAR Result	All four substances have a very high log KOW that is above the screening criterion for bioaccumulation in the PBT-assessment. The substances behave similar. Also KOWWIN predicts log KOWs approximately 0.8-1.0 log units smaller than COSMOtherm. The values of KOWWIN are nearer to the available experimental values.
Rationale	Not B-Screening criteria according to ECHA Guidance R.11 is log KOW < 4.5
Reliability	Reliable

Conclusion with regard to the regulatory purpose

The log KOW-values for all four substances are high and therefore a high bioaccumulation potential is expected. This expectation is confirmed by the available experimental BCF-values. All four substances have log KOW-values in the same region. While there seems to be a systematic shift between the results there is no such shift observed for the relative order of the values.

ANNEX 5.1: QMBI KOWWIN

	Information	Literature references or Links	Remarks		
0 - General	0 - General				
Model name and version	KOWWIN 1.68	Meylan, W.M. and P.H. Howard. 1995. Atom/fragment contribution method for estimating octanol-water partition coefficients. J. Pharm. Sci. 84: 83-92.			
W.a.23: software package	EPISUITE Estimation Programs Interface Suite [™] for Microsoft® Windows, v4.10	http://www.epa.gov/oppt/exposure/pubs/episuite.htm			
1 - Definition of					
Endpoint [units] (w.a. species and other relevant information)	n-octanol/water partition coefficient given as a logarithmic value				
2 – Definition of					
Brief description of algorithm and/or link to full definition	Log KOW = Σ(fi*ni)+ Σ (cj*nj) + 0.229	See Online help of KOWWIN	Derived by multiple regression of training set in a two step procedure: 1. Derivation of fi 2. Introduction of cj		
List of employed descriptors with units	fi: coefficient for each atom or fragment i; ni: number of times fragment/atom i occurs; cj: coefficient for correction instance j; number of times a structure that leads to a correction instance occurs	See Online help of KOWWIN, Appendix D	There are 157 different atoms and fragments defined and 278 correction factors that are employed when certain chemical classes or functional groups are present in the molecule for which an estimation is made		
Number of Chemicals in Training Set and Brief	2447 chemicals with measured log KOW-values from the PhysProp Database		Training Set Estimation Error: within $\leq 0.10 - 45.0\%$ within $\leq 0.20 - 72.5\%$		

²³w.a.: when applicable

description of it			within <= 0.40 - 92.4% within <= 0.50 - 96.4% within <= 0.60 - 98.2%
W.a.: Training set available at		List available at http://esc.syrres.com/interkow/KowwinData.htm	Within <= 0.00 - 96.2%
	the Applicability Domain		
W.a.: Definition of the Applicability Domain	Currently there is no universally		With exceedingly high or low log KOW the experimental errors for determination of log KOW will become larger and therefore the uncertainty. In such cases the predicted values will be more uncertain as well.
Limits of the	18.02 to 719.92 [g/Mol], for		
Applicability Domain	Structural Domain see Training Set		
	on the Validation of the Model		
Validation Set			
Туре	from different sources		
W.a.: Validation available at		List available at http://esc.syrres.com/interkow/KowwinData.htm	
Statistical information on validity	Validation Set Estimation Error: within <= $0.20 - 39.6\%$ within <= $0.40 - 66.0\%$ within <= $0.50 - 75.6\%$ within <= $0.60 - 82.5\%$ within <= $0.80 - 91.6\%$ within <= $1.00 - 95.6\%$ within <= $1.20 - 97.7\%$ within <= $1.50 - 99.1\%$ Interpretation of the model		Details available in Online help of KOWWIN

W.a.: Mechanistic basis of model	Fragment coefficients and correction factors reflect the impact of certain chemical fragments or functional groups
	on lipophilicity and thus on the log KOW.

ANNEX 5.2: QMBI COSMOtherm KOW

	Information	Literature references or Links	Remarks
0 - General			
Model name and version	COSMOtherm v C30_1201		The COSMOtherm model allows in principle the calculation of all partition properties of molecules. In this QMBI only the calculation of the KOW will be addressed
W.a.24: software package	COSMOtherm		
1 - Definition of	of Endpoint		
Endpoint [units] (w.a. species and other relevant information)	-		
2 – Definition of	of Algorithm		
Brief description of algorithm and/or link to full definition	$\begin{array}{l} \log & \text{KOW} & (\text{T}) = \\ \int p^{i}(\sigma)(\propto_{water}(\sigma;\text{T}) & - \\ \approx_{octanol}(\sigma;\text{T}) &)d\sigma & + \\ \approx_{i}^{C}(water, & \text{T}) & - \\ \approx_{i}^{c}(\text{octanol},; & \text{T}), \text{ where} \\ \approx_{i}^{c}(\text{S},\text{T}) = \text{RT}^{*} [\lambda_{0}^{*} \ln r_{i} \\ + \lambda_{1}^{*}(1 - (r_{i}/\underline{r} - \ln \underline{r}) + \\ \lambda_{2}^{*}(1 - q_{i}/\underline{q} - \ln \underline{q})] \text{ and } \underline{r} = \\ \Sigma_{I} x_{i}^{*}r_{i} \text{ and } \underline{q} = \Sigma_{i} x_{i}^{*}q_{i} \end{array}$	"COSMO-RS: From Quantum Chemistry to Fluid Phase Thermodynamics and Drug Design", Andreas Klamt, Elsevier Science Ltd., Amsterdam, The Netherlands (2005), ISBN: 0- 444-51994-7.	COSMOtherm implements the COSMO-RS theory. This theory interprets the interaction of molecules as an interaction of a larger ensemble of molecular surfaces calculated with Quantum Mechanical methods. Due to a treatment with statistical thermodynamics the macroscopic properties of interacting molecules like partition coefficients

²⁴w.a.: when applicable

		become available.
List of		
employed	[kcal/(mol K)], T:	
descriptors	temperature [K]; σ :	
with units	Screening charge density	
	or polarity, i.e. the	
	electrostatic screening of	
	a solute molecule by its	
	surrounding and its back	
	polarization in a region	
	with radius of ca. 0.5 Å ;	
	$p^{i}(\sigma)$: sigma profile of	
	molecule i, i.e. the sum of	
	the probability	
	distributions of all	
	possible σ ; $\propto_{water}(\sigma;T)$:	
	sigma potential of water at	
	temperature T, a sigma	
	potential can be interpreted	
	as the affinity of a molecule	
	for a surface of polarity σ ;	
	∝ _{octanol} (σ;T): sigma	
	potential of octanol at	
	temperature T; $\propto_i^c(S;T)$:	
	combinatorial contribution	
	to the chemical potential	
	of molecule i in solvent S	
	at temperature T; λ_0 , λ_1 ,	
	λ_2 : adjustable parameters,	
	ri: molecular volume of	
	substance i, qi: molecular	
	area of substance i, r:	
	overall volume of the	
	mixture, q: overall area of	
	the mixture.	
Number of	Original parameterisation:	While the principle theory is

Chemicals in Training Set and brief description of it	225 small- and medium- sized organic compounds with H, C, O, N, Cl atoms. The fitting was done for 650 experimental room- temperature parameters (Δ Ghydr, log(vapor pressure), log Koctanol- water, log Khexane-water, log Kbenzene-water, log Kdiethyl ether-water		applicable for all elements, the practical implementation needs some specific parameters to the QM-method used and the elements of the substance in question like the employed ratio for scaling the bonds of the QM- method and the van der Waals-coefficients
W.a.: Training set available at		"Refinement and Parametrization of COSMO-RS", Andreas Klamt, Volker Jonas, Thorsten Bürger and John C. W. Lohrenz, J. Phys. Chem. A 102, 5074-5085 (1998).	Since the original parameterization was done further adjustments were made and parameters for further elements were introduced. While the parameters are available in the software, to our knowledge the details of the new parameterisations were not disclosed
3 – Definition of	of the Applicability Domain		
W.a.:	There is no formal definition		
Definition of	of the applicability domain		
the Applicability Domain			
Limits of the	In principle the method is		
Applicability Domain	completely based on first- principles meaning there is no limit of the Applicability Domain.		
4 – Information	n on the Validation of the Model		
Validation Set Type	To our knowledge there is no single validation set but there are several citations in literature on the accuracy/validity of the		

	model		
W.a.:		Overview over publications:	
Validation		http://www.cosmologic.de/index.php?cosId=4150&crId=10	
available at			
Statistical	-		
information			
on validity			
5 – Mechanistic	Interpretation of the model		
W.a.:	The interaction of a solute		
Mechanistic	and a solvent is calculated in		
basis of	terms of a chemical		
model	potential. The difference of		
	the chemical potentials of		
	the solute in two different		
	solvents is the mechanistic		
	reason for partition effects.		

ANNEX 5.3: Analysis of QSAR prediction for UV-320 , UV-327, UV-328, UV-350

QSAR Model: KOWWIN and COSMOtherm (KOW)

Overall performance

	Result		Further description
Endpoint results	KOWWIN	UV-320: 6.27	All log KOW-values are high and in a
[unit]		UV-327: 6.91	similar region. There seems to be a
		UV-328: 7.25	systematic shift between the two
		UV-350: 6.31	models where KOWWIN predicts in
	COSMO-	UV-320: 7.39	general lower values.
	therm (KOW)		
		UV-327: 7.91	
		UV-328: 7.89	
		UV-350: 7.11	
Applicability domain	Yes		The molecules are in the range of all
			descriptors employed in the models
			and in the range of the molecular
			weight of the molecules in the
			training set of KOWWIN.
Similarity with	Yes		All fragments or elements of the
trainings set			molecules are represented in the
			Training Set of KOWWIN.
			COSMOtherm has no training set but
			is generally applicable.
Similar substances	Yes		See table next side
Model performance		e range of values	Experimental Values and predictions
for similar good, but absolute values seem		show a systematic shift but caution	
substances	to be slightly o	verestimated	has to be advised as the
			experimental values were not
			validated.
Other uncertainties	No		-

Overall conclusion	Reliable
Rational	As the models are applicable and results for similar molecules and two of the four models at hand show values in the same range it can be expected that the range is correctly predicted.

	Substance 1
Structure	
CAS-Nr.	70321-86-7
EU-Nr.	274-570-6
(Trade-)Name	UV-234
Descriptor value	KOWWIN : log KOW = 7.67 COSMOtherm: log KOW = 8.30
Predicted endpoint	See above
Experimental endpoint	> 6.5
Statistical performance	-

Results for similar substances

Rationale for the selection of similar substances

Substances 1 is structurally similar as it is a phenolic benzotriazole as the target molecule. It also has a sterical demanding side chain in ortho- and one in para-position to the hydroxyl group. The difference lies in the substitution of a phenyl group for a methyl group. Therefore is is probably to some degree more lipophilic as UV-327.

ANNEX 6: Monitoring Study Results for UV-320, UV-327, UV-328, UV- 350

Monitoring of phenolic benzotriazoles

Monitoring studies are summarized concerning the following phenolic benzotriazoles: UV-234 (CAS 70-321-86-7), -320 (CAS 3846-71-7), -326 (CAS 3896-11-5), -327 (CAS 3864-99-1), -328 (CAS 25973-55-1), -329 (CAS 3147-75-9), -350 (CAS 36437-37-3), -360 (CAS 103597-45-1) and -571 (CAS 125304-04-3). No monitoring studies were found for UV-928 (CAS 73936-91-1).

European studies:

Brorström-Lundén et al. (Brorström-Lundén et al., 2011) published a screening study on benzotriazoles (UV-234, -320, -327, -328, -329, -360). Phenolic benzotriazoles may to a large extent enter Sweden through imported finished goods. Emissions via diffuse sources were assumed as the main pathway of benzotriazole UV-absorbers to the environment. The sampling program was therefore focused on emissions in urban environments (Stockholm area and smaller city Borås). In addition background sites were included and two sites with potential point sources. Benzotriazoles were analyzed using an LC-MS system including a tandem mass-spectrometer. Detection limits vary with analyzed substance and sample. Compared to other studies the detection limits for sediment, soil, particles, WWTP sludge and fish are high.

Compartment	Detection limits	Compartment	Detection limits
Air	0.01 – 0.48 ng/m ³	storm water	0.03 – 0.1 ng/L
air deposition	30 – 200 ng/m ² day	landfill effluent particles	0.7 -1.6 µg/g dw
surface water	0.03 – 0.09 ng/L	landfill effluent	0.08 – 0.5 ng/L
Sediment	0.2 – 12 µg/g dw	WWTP effluent particles	61 – 130 µg/g dw
Soil	0.1 – 0.9 µg/g dw	WWTP effluent	0.04 - 0.1 ng/L
Fish	0.3 – 1.9 µg/g dw	sludge	0.1 – 0.6 µg/g dw

Table 22: Detection limits in the investigation of Brorström-Lundén et al.

In air samples 4 benzotriazole UV-absorbers were detected (UV-320, -327, -329, -360). Concentrations were similar in background and urban air. However, the highest concentration was measured in Stockholm. Only two compounds were detected in atmospheric deposition (UV-327, -329). The deposition was higher at the urban site.

Table 23: Concentrations of phenolic benzotriazoles in air and atmospheric deposition in Sweden

Substance	Air		Deposition		
	detected in x of y samples [x/y]	concentration [ng/m ³]	detected in x of y samples [x/y]	deposition flux [ng/m² day]	
UV-234	0/8	-	0/4	-	
UV-320	3/8	0.024 - 0.67	0/4	-	
UV-327	6/8	0.40 - 25	3/4	<100-320	
UV-328	0/8	-	0/4	-	
UV-329	5/8	< 0.15 - 3.0	3/4	<100-331	
UV-360	1/8	0.40	0/4	-	

Several benzotriazoles were found in soil, in rather similar concentrations at the background and the urban locations (UV-320, -327, -328, -329). There were differences in the occurrence among the individual substances at the different locations. According to the authors the highest concentration of a single substance (UV-329) was found in Soil 500 m from a busy road in the Stockholm area. However, according to the annex of the study such a high concentration was also found for UV-327 in another urban sample. Since only 4 samples were analyzed altogether, the results should generally be interpreted with care.

Several of the benzotriazoles were frequently detected in surface water (UV-320, -327, -328. - 329). The concentrations were mostly similar at background and urban locations. In sediments

the distribution among different substances varied for the different sampling sites. Peaks of single substances occurred both at background and urban locations; the lower concentration levels were similar at different locations.

Three of the benzotriazoles were found in fish, both at urban and background locations (UV-324, -327, -329). The highest concentration was found at the background location (UV-327). The concentrations found in Swedish fish are 1000fold higher than those found in Japanese fish. The reason for this is unknown. The authors note however that most substances are not detected and the levels found are quite close to the detection limit of the method used.

Substance	Soil		Fish	
	detected in x of y	concentration	detected in x of y	concentration
	samples [x/y]	[µg/g dw]	samples [x/y]	[µg/g dw]
UV-234	0/4	-	1/4	0.26
UV-320	1/4	0.91	0/4	-
UV-327	3/4	0.66-3.7	3/4	2.3-9.8
UV-328	1/4	0.74	0/4	-
UV-329	3/4	0.79-3.7	3/4	1-2.5
UV-360	0/4	-	0/4	-

Table 24: Concentrations of phenolic benzotriazoles in soil and fish in Sweden

Table 25: Concentrations of phenolic benzotriazoles in surface water and sediment in Sweden

Substance	Surface water		sediment	
	detected in x of y	concentration	detected in x of y	concentration
	samples [x/y]	[ng/L]	samples [x/y]	[µg/g dw]
UV-234	0/6	-	0/6	-
UV-320	3/6	0.55-0.94	5/6	0.16-3
UV-327	4/6	0.11-0.39	6/6	1.6-35
UV-328	6/6	1.3-10	4/6	0.65-1.3
UV-329	6/6	0.25-2.4	4/6	0.81-33
UV-360	1/6	0.16	3/6	0.42-2.9

All benzotriazoles but UV-360 were detected in WWTP effluent and all substances were detected in sludge from WWTPs. However, there were differences both in concentration levels and in distribution among the different benzotriazoles between the WWTPs. A different distribution among the substances was also found in effluent and sludge. Only one sample of WWTP effluent particles was analyzed and only UV-327 was detected in this sample (270 μ g/g dw).

Table 26: Concentrations of phenolic benzotriazoles in WWTP effluent and sludge in Sweden

Substance	effluent WWTP		sludge WWTP	
	detected in x of y	concentration	detected in x of y	concentration
	samples [x/y]	[ng/L]	samples [x/y]	[µg/g dw]
UV-234	1/5	0.11	8/8	2.1-7.3
UV-320	1/5	4	6/8	0.84-2
UV-327	4/5	0.12-0.48	7/8	0.54-17
UV328	5/5	6.8-15	4/8	2.8-37
UV-329	5/5	0.87-4.9	7/8	2.3-15
UV-360	0/5	-	8/8	4.6-23

All substances but UV-360 were found in landfill leachates, all substances but UV-329 occurred in storm water. In one sample of landfill effluent particles UV-327, -328 and -329 were detected in concentrations of 4.3, 3.1 and 6.1 μ g/g dw, respectively.

Table 27: Concentrations of phenolic benzotriazoles in effluent landfill and storm water in Sweden

Substance	effluent landfill		storm water	
	detected in x of y samples [x/y]	concentration [ng/L]	detected in x of y samples [x/y]	concentration [ng/L]
UV-234	2/3	0.16 and 0.5	4/4	0.06-0.31
UV-320	2/3	7.3 and 23	1/4	0.73

UV-327	2/3	0.45 and 1.3	3/4	0.13-0.17
UV-328	3/3	7-91	3/4	0.19-1.3
UV-329	1/3	17	0/4	-
UV-360	0/3	-	2/4	0.17 and 0.28

In summary widespread occurrence of benzotriazoles in the Swedish environment was observed both in background and urban areas. The substances occurred in all environmental matrices included in the study: air, deposition, surface water, sediment, soil and biota. Diffuse spreading through WWTPs, landfills and storm water may be important for the occurrence in the environment. Levels measured in WWTP effluents and sludge indicate widespread diffusive sources via use of products. The benzotriazoles with the highest usage volume in Sweden (UV-327, UV-328) were also most often found in the highest concentrations.

The authors conclude that on a national scale air transport may be a significant source of the compounds and that the substances are stable enough to undergo atmospheric long range transport.

Carpinteiro et al. (Carpinteiro et al., 2010a) used headspace solid-phase microextraction followed by gas chromatography tandem mass spectrometry for the sensitive determination of benzotriazole UV-stabilizers in water samples (UV-326, -327, -328). The limit of quantification was < 2 ng/l. The developed methodology was used to investigate the presence of benzotriazoles in filtered river water (3 samples), two samples taken in the inlet and outlet streams of an urban WWTP and four additional specimens of raw wastewater provided by a local laboratory. Phenolic benzotriazoles were not detected in river water and treated wastewater. In raw wastewater samples UV-327 was not detected, whereas UV-326 and -328 were each found in 4 of 5 samples in concentrations ranging from 3.5-57 ng/L and 1-19 ng/L, respectively.

Carpinteiro et al. (Carpinteiro et al., 2010b) also investigated benzotriazole UV-stabilizers in indoor dust samples (UV-326, -327 and -328). Pressurized liquid extraction and gas chromatography followed by tandem in time mass spectrometry were used. The limits of quantification were between 4 and 9 ng/g. Procedural blanks showed small peaks at the retention time of some species. The source of this contamination may be related to the trend of target compounds to be retained on solid surfaces. Glass material, extraction cells and connections in the extraction system might contribute to the presence of benzotriazole UV-stabilizers in procedural blanks due to carry over problems.

Dust was collected with domestic vacuum cleaners equipped with paper filter bags from several private houses (5 samples), vehicle cabins (3 samples) and an administrative building (1 sample). It is not stated in which country the dust was collected. However, it is assumed that it was collected in Spain. The dust fraction < $60 \mu m$ was used for the study. In addition a house dust reference material from USA was acquired. This sample was used to confirm the ubiquity of benzotriazole UV-stabilizers in dust although no certified or indicative values of their levels in the reference material were available.

UV-326, -327 and -328 were found to be ubiquitous in dust, with measured values from 22 to >600 ng/g. Moreover, UV-326 was found in one car cabin dust sample at a concentration of almost 5 μ g/g.

	UV-326	UV-327	UV-328
private house 1	42	86	46
private house 2	58	101	127
private house 3	333	29	100
private house 4	73	22	68
private house 5	269	52	149
public building	676	131	62
car cabin 1	4880	48	88
car cabin 2	522	127	124
car cabin 3	170	43	52
US dust reference material	121	322	259
Min-Max (Mean) of all samples except US material	42 - 4883 (780)	22 - 127 (71)	46 - 149 (91)

Carpinteiro et al. (Carpinteiro et al., 2012b) combined stir-bar sorptive extraction and liquid desorption with large volume injection-gas chromatography-mass spectrometry for the determination of benzotriazole UV-stabilizers in wastewater matrices. UV-320, -326, -327 and -328 were measured in urban sewage waters. Grab samples of wastewater were obtained from inlet and outlet streams of two urban WWTPs, equipped with primary and activated sludge treatment units, located in Portugal and Spain. The limits of quantification were between 4 and 10 ng/L. Because of the existence of significant concentrations of phenolic benzotriazoles associated with dust particles it is highly recommended to protect laboratory material from deposition of particulate matter. The efficiency of the extraction is sample dependent; therefore, the standard addition method is required for the accurate quantification of the substances in wastewater matrices.

Table 29: Average concentrations of phenolic benzotriazoles in wastewater matrices (n = 3 replicates) [ng/L]

Place, date	type	UV-320	UV-326	UV-327	UV-328
Portugal,	raw wastewater	24	26	85	76
Nov. 2010	treated wastewater	n.d.	n.d.	31	21
Spain,	raw wastewater	n.d.	40 (6)	n.d.	53
Jan. 2011	treated wastewater	n.d.	n.d.	n.d.	n.d.
Spain,	raw wastewater	n.d.	34	22	65
Feb. 2011	treated wastewater	n.d.	n.d.	n.d.	n.d.

n.d. = not detected

Carpinteiro et al. (Carpinteiro et al., 2012a) also measured benzotriazole UV-absorbers in sediments. Matrix solid-phase dispersion followed by gas chromatography tandem mass spectrometry was used. The limit of quantification of the method was 3 ng/g for UV-320, -326, -327 and -328. Ten samples of river and estuarine sediments with different carbon contents were investigated. Fresh sediment samples were air-dried in the hood for several days then sieved. The fraction with the particle size < 0.3 mm was considered in the study. In 6 of the 10 sediment samples quantifiable levels of UV-absorbers were detected:

Table 30: Concentrations of benzotriazole UV-absorber species measured in sediment samples (paricle fraction < 0.3 mm, n=3 replicates, - = not detected)

Sample	total carbon	UV-320	UV-326	UV-327	UV-328
	[%]	[ng/g]	[ng/g]	[ng/g]	[ng/g]
1	3.0	5.6	32	15	56
2	3.9	-	-	10.3	10
3	5.5	-	7.8	-	8.3
4	4.6	-	-	9.5	11.2
5	2.2	-	-	-	7.9
6	8.0	-	15	-	8

Unfortunately the origin of the sediment samples is not mentioned in the study. According to the acknowledgements some of the analyzed sediment samples were supplied by the German Federal Institute of Hydrology. However, the authors could not specify which samples were from Spain and which were from Germany (personal communication April 2012).

Montesdeoca-Esponda et al. (Montesdeoca-Esponda et al., 2012) used on-line solid-phase extraction coupled to ultra-performance liquid chromatography with tandem mass spectrometry detection (SPE-UPLC-MS/MS) for the determination of UV-326, -327, -328, -329, -360 and -571 in samples from WWTP effluents and coastal marine water from Spain. The detection limits and quantification limits achieved were in the range of 0.6-4.1 ng/L and 2.1-14 ng/L. The analytical method allowed simultaneous determination of the compounds in liquid samples with satisfactory recoveries and reproducibility, except for UV-360, which cannot be completely eluted from the cartridge due to its high octanol-water partition coefficient and molecular mass.

Seawater samples were collected from six beaches around the Gran Canaria Island in Spain (2 samples per beach), wastewater samples were collected from seven WWTPs of Gran Canaria

Island. All substances studied were detected in the wastewater samples (see table). In seawater samples only UV-360 was found (6 of 12 samples, 3.6 - 5.2 ng/L).

	detection frequency	concentration(s) [ng/L]
UV-326	1/7	11
UV-327	1/7	4.8
UV-328	5/7	6.2 - 13
UV-329	1/7	4.0
UV-360	2/7	5.9 and 6.6
UV-571	0/7	not detected

Table 31: Concentrations of phenolic benzotriazole UV-absorbers in samples of WWTP effluents of Gran Canaria Island

Soil and suspended solids samples from the German Environmental Specimen Bank were analyzed for UV-234, -320, -326, -327, -328, -329 and -350 at the University of Santiago de Compostela (Rodríguez Pereiro and Casado Agrelo, 2012). Samples were extracted using the matrix solid-phase dispersion (MSDP) technique, with an integrated clean-up step. A GC-MS/MS method was used with a hybrid quadrupole time-of-flight mass spectrometer furnished with an electronic impact source. The limits of quantification were 2 ng/g per compound.

Samples were from sites with high anthropogenic influence and from background sites. Five soil samples taken in 2010 and five samples of suspended particulate matter taken in 2011 were analyzed. Soil samples were 3 litter samples, one root network sample and one top soil sample. All soil samples revealed target compound levels below the limits of quantification, also for the soils from Saarbruecken-Staden (root network) and Duebener Heide/Leipzig (litter, top soil) which are assumed to be more anthropogenically influenced. Concentrations of phenolic benzotriazoles in suspended solids samples are shown in Table 32.

Table 32: Concentrations of phenolic benzotriazoles in suspended solids samples from Germany

Suspended solids sample	UV-234 [ng/g dw]	UV-320 [ng/g dw]	UV-326 [ng/g dw]	UV-327 [ng/g dw]	UV-328 [ng/g dw]	UV-329 [ng/g dw]	UV-350 [ng/g dw]
Danube / Jochenstein	n.d.						
Rhine /Weil	n.d.	n.d.	26	n.d.	26	n.d.	n.d.
Elbe / Cumlosen	8.1	n.d.	4.6	n.d.	n.d.	n.d.	n.d.
Saale / Wettin	15	n.d.	17	n.d.	n.d.	n.d.	n.d.
Saar / Rehlingen	17	n.d.	17	n.d.	n.d.	2.0	n.d.

n.d. = not detected

Suspendid solids from the river Elbe and its tributary Saale showed similar patterns, with higher levels for the tributary Saale. Patterns for suspended solids from the rivers Saale and Saar are comparable. Both rivers revealed high burdens also for other substances. The Rhine site Weil downstream Basel is influenced by the Swiss chemical industry and has a different pattern (higher level of UV 326, only site with UV 328). The Danube site at Jochenstein was selected because of low burdens and displayed levels below the limits of quantification.

Japanese studies:

Nakata et al (Nakata et al., 2009a) studied occurrence and concentrations of UV-320, -326, -327 and -328 in marine organisms and sediments from the Ariake Sea, western Japan. 16 coastal and river sediments were collected during 2006-2007. Five of the sediment samples were taken in a heavily polluted river. 55 biota samples were collected during 2004 and 2007:

- tidal flat organisms: lugworm, lamp shell, oyster, clam, gastropod, crustaceans (crab, shrimp), fishes (herbivorous and omnivorous mudskippers)
- shallow water species: crustaceans (crab, shrimp), teleost fish (flathead, solefish, right eye flounder, sandperch, sweetlips, mullet, sea bass, hairtail), cartilaginous fish (eagle ray, hammerhead shark)

• coastal birds (spot-billed duck, mallard).

Depending on the species, the whole body, soft tissue, hepatopancreas and liver samples were analyzed. 16 coastal and river sediments were also collected around the Ariake Sea during 2006-2007. UV-stabilizers were detected in all biota and sediment samples. In biota UV-326, - 327 and -328 were the dominant compounds at levels of 0.1-55 ng/g ww. Concentrations of UV-320 in samples were low, it could be detected only in tidal flat organisms and some shallow water species. This may be due to small amounts of use of this compound in Japan since its domestic production and use have been restricted.

In general, concentrations of UV-stabilizers in tidal flat organisms were greater than those in shallow water species. The average concentrations of UV-320 and UV-326 in tidal flat species were approximately 10- to 20-fold higher than those in shallow water organisms. The tidal flat clam showed the highest concentrations of UV-320 and UV-326 at 74 ng/g and 219 ng/g (lw) respectively. Elevated concentrations of UV-326 were also found in oysters and gastropods in tidal flat area. These results imply the presence of phenolic benzotriazoles in sediment, resulting in accumulation of these compounds in benthic organisms. The low concentrations of UV-326 in shallow water species might be explained by low BCF of this compound, as compared with other benzotriazole UV-filters. In addition the authors speculate that biodegradation of UV-326 in shallow water organisms may be a possible reason for low accumulation of this compound.

UV-327 was most frequently detected in the organisms investigated. The average concentrations of UV-327 in tidal flat organisms were only 2-fold higher than those in shallow water species. The tidal flat clam, crab and herbivorous mudskipper contained high concentrations of UV-327 (> 100 ng/g lw), followed by gastropods and oysters. In shallow water fishes such as mullet, sea bass and young sea bass, concentrations of UV-327 were 3- to 4-fold higher in liver than in carcass. These results are consistent with the concentration profiles of UV-328 in mullet, suggesting the preferential accumulation and less biodegradation of this compound in the liver of some fish species. Omnivorous birds accumulate UV-327 in the liver, at average concentrations of 90 ng/g (lw) in a spot-billed duck and 59 ng/g in mallards. This suggests bioaccumulation in higher trophic species in the aquatic food chain.

Concentrations of UV-328 in biota were variable and species-specific. The highest concentration was found in tidal flat gastropod at 460 ng/g (lw), followed by mullet (120 ng/g lw in whole body and 250 ng/g lw in liver) and hammerhead shark (130 ng/g lw in liver) collected from shallow waters. The oysters and clams in tidal flat contained high concentrations of UV-328, at >100 ng/g lw. The large variations in UV-328 concentrations observed in this study might be due to differences in retention and metabolism of this compound in marine organisms.

As described above, the concentrations of benzotriazole UV-stabilizers in tidal flat organisms were higher than those in shallow water species. In addition, clams, oysters and gastropods presented high concentrations of UV-320, UV-326 and UV-328 rather than crabs and fishes, although the former species are at lower trophic levels in the tidal flat ecosystems. There is no positive correlation between the concentrations and the trophic status of organisms in marine ecosystems.

The benzotriazole UV-stabilizers were detected in 11 coastal sediments analyzed, at total concentrations of several ng/g dw. UV-328 was found at the highest concentrations (average $6.4 \pm 4.0 \text{ ng/g dw}$), followed by UV-326 ($3.7 \pm 3.0 \text{ ng/g dw}$), UV-327 ($3.2 \pm 2.6 \text{ ng/g dw}$) and UV-320 ($0.9 \pm 0.6 \text{ ng/g dw}$). The composition of the UV-stabilizers among the sediment samples was less variable than in biota. Extremely high concentrations were found in five sediments from the highly polluted Omuta River. Highest concentrations of UV-320, -326, -327 and -328 reached 14, 200, 190 and 320 ng/g dw, respectively. Significant correlations were found in sediment concentrations between UV-326 and 327, UV-326 and 328, and UV-327 and 328 in the Ariake Sea. Significant correlations were also found between UV-stabilizer concentrations and organic carbon contents in sediment.

Table 33: Concentrations of benzotriazole UV-stabilizers in tidal flat and shallow water organisms collected in Japan

	UV-320	UV-326	UV-327	UV-328
	[ng/g ww]	[ng/g ww]	[ng/g ww]	[ng/g ww]
10 tidal flat organisms	< 0.05 - 0.60	< 0.10 - 2.5	< 0.12 - 3.6	0.35 - 14

10 marine shallow water organisms	< 0.05 - 0.09	< 0.10 - 0.32	< 0.12 - 2.3	0.19 - 8.7
6 marine shallow water organisms (liver)	< 0.05 - 7.0	< 0.10 - 5.6	2.4 - 13	< 0.15 - 55
2 species of water fowl (liver)	< 0.05	< 0.10	2.6 3.4	< 0.15

Table 34: Concentrations of benzotriazole UV-stabilizers in sediments in Japan

	UV-320 [ng/g dw]	UV-326 [ng/g dw]	UV-327 [ng/g dw]	UV-328 [ng/g dw]
marine and estuarine sediments $(n = 11)$	0.3 - 2.3	1.5 - 12	1.6 - 9.9	7.9 - 40
Omuta River sediments $(n = 5)$	2.6 - 14	23 – 200	16 - 190	18 - 320

Nakata et al. (Nakata et al., 2009b) also investigated occurrence and concentrations of UV-320, 326, 327 and 328 in marine organisms collected from the Ariake Sea, western Japan. 51 marine organisms, such as lugworms, mussels, oysters, crustaceans, fish, birds and marine mammals were collected during 2001 and 2005. 12 sediments were collected from the same region in 2007. Analyses were done via GC-MS.

UV-filters were detected in most marine organisms in the study. Highest concentrations were found in lower benthic organisms, gastropods, collected from the tidal flat area (UV-328 > 400 ng/g lw). UV-328 and -326 were the dominant components in these organisms. In shallow water species, elevated levels were found in the liver of mullet, a benthic fish (UV-328 > 200 ng/g lw). Higher trophic species, such as sharks, marine mammals and birds accumulate organic UV-filters. UV-328 and -327 were dominant in finless porpoises and mallards, respectively. The results suggest significant bioaccumulation of UV-filters through the marine food-webs.

The substances were also detected in surface sediments from the Ariake Sea (average concentration: several ng/g dw). High concentrations of UV-filters were found in the Omuta River sediments, at levels ranging from 2.3-320 ng/g dw. Significant correlations were found between concentrations and organic carbon contents in sediments. No more details are given.

In order to understand the geographical distribution of UV-filters, blue and green mussels from 10 Asian countries and regions were collected during 1998 and 2005 and analyzed (Cambodia, China, Hong Kong, India, Indonesia, Japan, Korea, Malaysia, the Philippines, Vietnam). Only qualitative information is given on this investigation. UV-filters were detected in most mussel samples, indicating the widespread use of these compounds in Asian coastal regions. In general, UV-326 was the dominant compound, whereas UV-320 was detected only in several samples collected from Japan. The UV-filters concentrations were high in mussels from Korea, Japan and Hong Kong. Low residue levels of UV-filters were found in samples from India and Vietnam. These results suggest different usage values of UV-filters among countries and regions in Asia. Concentrations in mussels showed great spatial variations in Korea and Japan, which may be due to the distance between the sampling points and the sources of UV-filters, such as WWTPs. Significant positive correlation was determined in concentrations between UV-327 and UV-328 in mussels.

Nakata and Shinohara (Nakata and Shinohara, 2010) analyzed UV-320, -326, -327 and -328 in influent, effluent and sewage sludge samples collected from 5 WWTPs located in a town (population 680,000) in Japan. Samples were taken in May and October 2009. The wastewater flows were 140,000, 29,300, 9,300, 53,300 and 63,200 m³/d, respectively. The treatment process included activated sludge method in all WWTPs. In the biggest WWTP (East WWTP) influent samples were collected at 9:00, 12:00, 15:00, 18:00 and 21:00 (n = 5), to study time-dependent variations of target substance concentrations. Influent and effluent samples were also obtained from the 4 other WWTPs (n = 1 / sample). Two sewage sludge samples were also collected from each of the five WWTPs (n = 10). The detection limits ranged from 2.1 to 8.7 ng/L in this study (limits of quantification not given).

Benzotriazole UV-stabilizers were detected in all influents collected from East WWTP at every three hours during 9:00 to 21:00. UV-326 showed the highest concentrations in influents, followed by UV-328 and -327.

Time of sampling	9:00	12:00	15:00	18:00	21:00	Average ± standard deviation
UV-326	26	24	23	19	28	24 ± 3.7
UV-327	17	11	10	20	5.6	12 ± 5.6
UV-328	23	20	17	14	15	18 ± 3.9

Table 35: Concentrations [ng/L] of benzotriazole UV-stabilizers in influents of East WWTP

Concentration in	UV-326	UV-327	UV-328
influent (9 samples) [ng/L]	24 - 78	< 8.7 - 12	18 - 52
effluent (5 samples) [ng/L]	3.0 - 4.5	< 8.7	2.1 - 2.9
sludge (10 samples) [ng/g dw]	760 - 1800	120 - 200	430 - 570

Benzotriazole UV-stabilizers were detected in most samples analyzed and UV-326 was the dominant compound in influents (mean: 46 ng/L), followed by UV-328 (34 ng/L). UV-327 was detected in two influents at concentrations of 9.2 and 12 ng/L. UV-320 was not identified in any of the samples, probably because its domestic production and use have been restricted in Japan. These results imply a large amount of production and usage of UV-326 compared with other benzotriazole UV-stabilizers in Japan. Concentrations in the effluents were generally < 5 ng/L, suggesting an elimination of these compounds during wastewater treatment. The removal rates of UV-326 and -328 were >90% in the effluents, but high concentrations of benzotriazole UV-stabilizers were detected in sewage sludge samples of WWTPs, at high levels indicating adsorption to organic carbon in sewage sludge. The mean carbon percentage of sewage sludges was 31 \pm 2.2 %. Partition coefficients (Kp) were calculated at a moisture content of 80% in sludges. The values are 7,200 \pm 3,900 L/kg for UV-326 and 4,200 \pm 970 L/kg for UV-328.

Nakata et al. (Nakata et al., 2010) also detected benzotriazole UV-stabilizers in the blubber of marine mammals. They analyzed UV-320, -327 and -328 in finless porpoises (Neophocaena phocaenoides) collected from the Yatsushiro Sea, Ariake Sea and Tachibana Bay, Japan, in 1999, 2008 and 2009, respectively. All animals were stranded or accidentally caught by fishing net. Detection limits were 0.05, 0.12, 0.15 ng/g for UV-320, -327 and -328, respectively.

Table 37: Concentrations of benzotriazole UV-stabilizers [ng/g ww] in the blubber of finless porpoises

sample no.	1	2	3	4	5
sampling year	1999	1999	2008	2009	2009
lipid content [%]	81	83	87	59	91
UV-327	4.5	9.5	6.3	31	18
UV-328	20	64	11	34	16

UV-320 was not detected in the samples, which is attributed to its restriction in Japan in 2007. The mean concentrations and standard deviations of UV-327 and UV-328 in five blubber samples were 19 \pm 19 ng/g lw and 38 \pm 28 ng/g, respectively, reflecting the higher consumption of UV-328 in Japan.

The authors cite a study showing a high concentration of UV-327 in the liver of a common cormorant (220 ng/g) collected from Hokkaido, northern Japan (respective reference in Japanese). While the concentrations of UV-327 in finless porpoises were lower than those in seabirds, the occurrence of UV-327 in marine mammals suggests the potential bioaccumulation in higher trophic species through the aquatic food chain.

According to the authors it has been reported that UV-327 concentrations in seawater from four coastal areas of Tokyo Bay were less than 0.5 ng/L and that the geometric mean concentration in river, lake and coastal water samples (n = 44) was 0.12 ng/L (respective references in Japanese). On the basis of these water concentrations the BAF of UV-327 between water and finless porpoises was estimated to be 33,300. Applying the same water concentrations to the calculation of a BAF of UV-327 in small fish inhabiting the same regions

results in a value of 3250, which is comparable to the values found under laboratory conditions (3400 to 9000).

UV-328 was not detected in the liver of seabirds, although UV-327 was present in the samples (Nakata et al. 2009b). The log Kow of UV-328 is the highest (8.28 reported in study) among the analyzed substances, but the BCF in fish was relatively low, 570-1400 and 620-2700 at the exposure concentrations of 0.1, 0.01 for 60 day, respectively (respective reference in Japanese). However, UV-328 showed a very high BCF, 36,000, between water and innards of fish (respective reference in Japanese). The authors conclude that the bioaccumulation profiles of UV-328 in marine organisms might be related to different retention and metabolism of this compound among species. The occurrence of UV-328 in finless porpoise may imply a low potential for biotransformation of this compound in this species. Finally it is stated that benzotriazole UV-stabilizers appear to be persistent and bioaccumulative in the aquatic food chain.

Kameda et al. (Kameda et al., 2011) measured 18 sun-blocking agents, among them UV-234, -326, -327, -328 and -329 in water and sediment collected from 22 rivers, 4 WWTP effluents and 3 lakes in August and September 2008 in Japan. Phenolic benzotriazoles are the most widely used UV-light stabilizers in Japan. WWTP sediment samples were collected from the river at the point of WWTP effluent discharge. In order to estimate contribution of sun-blocking agents from domestic wastewater to those in surface water and sediment, an indicator chemical for domestic wastewaters and WWTP effluents was also measured (HHCB = 1,3,4,6,7,8-Hexahydro-4,6,6,7,8,8-hexamethyl-cyclopenta-[g]-2-benzo-pyrane, a polycyclic musk, CAS 1222-05-5). The sampling sites represent 5 different groups:

- 2 streams with direct inputs of domestic wastewater (S1,S2)
- 4 WWTP effluents (ST1-ST4), conventional activated sludge treatment plants,
- 6 rivers heavily polluted by industrial and domestic wastewaters (H1-H6),
- 12 moderately contaminated rivers (M1-M12),
- 2 little rivers and 3 lakes as background sites (BG1-BG5).

Background sites did not receive domestic or industrial wastewater, but have possible slight sources (atmosphere deposition, recreational activities). In spite of considerable care, UV-328 was detected in blank samples. According to the authors this contamination was caused by analyte in indoor floor dust in the laboratory during experiments. The measured concentrations were corrected by the use of blanks upon each analysis. The limits of detection ranged from 0.1 ng/l to 3.0 ng/l and from 0.05 ng/g dw to 1.0 ng/g dw except for UV-328 which had a LOD of 10 ng/g dw.

The profiles of sun-blocking agents in surface water demonstrated site-specific differences at each sampling site. UV-328 was one of the dominant sun-blocking agents measured in water samples from heavily and moderately polluted rivers. The maximum level of UV-328 in heavily polluted rivers was near the lowest chronic NOEC of the substance estimated by EPI Suite (7 μ g/L). UV-234 and UV-329 were neither detected in water samples from surface waters nor from WWTP effluents. At the background sites none of the phenolic benzotriazoles analyzed were found in water samples.

analyte		UV-326	UV-327	UV-328
streams (S1, S2)	Occurrence	1/2	1/2	1/2
	mean detected a [ng/L]	16	5	70
	range [ng/L]			
WWTP effluents (ST1-ST4)	Occurrence	1/4	1/4	3/4
	mean detected [ng/L]	13	2	62
	range [ng/L]			47-88
heavily polluted rivers (H1-H6)	Occurrence	1/6	1/6	4/6
	mean detected [ng/L]	9	1	701
	range [ng/L]			149-4780
moderately polluted rivers	Occurrence	5/12	6/12	8/12
(M1-M12)	mean detected [ng/L]	2	1	152
	range [ng/L]	1-22	1-6	30-583

Table 38: Concentrations of phenolic benzotriazoles in water samples. UV-234 and 329 were not detected.

background sites (BG1-BG5)	Occurrence	0/5	0/5	0/5
	mean detected [ng/L]			
	range [ng/L]			

a geometric mean calculated from detected samples

 Table 39: Concentrations of phenolic benzotriazoles in sediment samples

analyte		UV-234	UV-326	UV-327	UV-328	UV-329
streams	Occurrence	1/2	2/2	2/2	2/2	1/2
(S1, S2)	mean detected a [µg/kgb]	1266	7.8	4.7	102	16
	range [µg/kgb]		0.1-110	0.6-37	10-1146	
WWTP	Occurrence	0/4	4/4	4/4	3/4	0/4
effluents	mean detected [µg/kg]		0.8	0.5	13	
(ST1-ST4)	range [µg/kg]		0.4-5.4	0.3-1.0	10-85	
heavily	Occurrence	4/6	5/6	5/6	6/6	3/6
polluted rivers	mean detected [µg/kg]	99	4.7	2.4	117	26
(H1-H6)	range [µg/kg]	38-324	0.9-45	0.7-18	21-1735	7.4-269
moderately	Occurrence	8/12	12/12	10/12	9/12	3/12
polluted rivers	mean detected [µg/kg]	47	1.8	0.9	59	0.6
(M1-M12)	range [µg/kg]	18-315	1.0-5.0	0.4-2.6	10-213	0.1-4.3
background	Occurrence	3/5	2/5	2/5	3/5	0/5
sites	mean detected [µg/kg]	39	1.2	0.7	58	
(BG1-BG5)	range [µg/kg]	8.3-113	1.1-1.3	0.5-1.1	29-89	

a geometric mean calculated from detected samples

b µg/kg dw

UV-234, -326, -327 and -328 were detected in most sediments. The compositions of sunblocking agents in sediment were quite similar among the five sampling site groups. The highest geometric mean concentrations of 18 sun-blocking agents in sediments were detected in streams and in heavily polluted rivers. The highest contributions to the total concentrations were those of UV-234 and -328. These two substances accounted for 70-80% of the total contaminants identified at all sediment sampling sites.

The results demonstrate that high concentrations of phenolic benzotriazoles were accumulated in sediment receiving not only chemical plants effluent, but also residential wastewaters, WWTP effluent and surface runoff.

UV-234, -326, -327 and -328 were significantly correlated with HHCB in sediments from rivers and lakes. According to the authors this shows that a large input of these substances is from domestic wastewater or WWTPs. It also suggests that their behavior in rivers and lakes, such as partitioning and attenuation, is similar to that of HHCB. UV-329 had no significant correlation with HHCB in sediments.

UV-326 had a strong linear correlation between UV-327 as well as UV-328 in all sediments. Since UV-stabilizers are often used as mixtures, the ratios observed in sediments may reflect their compositions in the products. The authors suggest that their (degradation) behavior may be also quite similar.

In a presentation Nakata (Nakata, 2011) showed graphs with concentrations of UV-326, -327 and -328 in mussels from 10 Asian countries and in mussels from the USA mussel watch program. All data cited are taken from the graphs. 45 samples were taken during 2003 and 2005.

UV-326 was detected in mussels from 7 of the 10 Asian countries. Highest concentrations were detected in mussels from Japan and Korea (ca. 1.5 and ca. 1.2 μ g/g lw, respectively). UV-327 was detected in 6 of the 10 countries with highest concentrations in Hong Kong and Korea (ca. 0.3 μ g/g lw). UV-328 was detected in 8 of the 10 countries with highest concentrations in Hong Kong and Korea (ca. 0.8 μ g/g lw).

In the USA samples were taken from blue mussels at 17 locations (n = 34) on the west coast (Alaska, Oregon, California) in 1994/95 and 2004/05. UV-326 and -327 were detected in most samples (14/17). Concentrations of UV-326 were similar to those measured in Japan and Korea. However, the maximum concentration was lower (ca. 0.7 μ g/g lw). Concentrations of UV-327 were higher than in Japan, but slightly lower than in Korea and had a maximum of ca. 0.25 μ g/g lw. UV-328 was detected in few samples, only, and showed a maximum of ca. 0.3

µg/g lw.

In an article Nakata et al. (Nakata et al., 2012) published more details on the mussel analyses. However, some more samples were included and other samples were excluded, so the results published in the article differ somewhat from those given in the presentation. Compounds analyzed were UV-320, -326, -327 and -328. 53 samples of blue and green mussels were collected from Cambodia, China, Hong Kong, India, Indonesia, Japan, Korea, Malaysia, Philippines and Vietnam during 2003 and 2007. In addition the analysis comprised 15 samples of blue mussels from the Pacific coast of the USA collected during 2004 and 2005. Liquid extraction and GC-MS in selective ion monitoring (SIM) mode was used. The limits of detection are given as 0.05, 0.1, 0.12 and 0.15 ng/g ww for UV-320, -326, -327 and -328, respectively.

Table 40: Mean concentrations of phenolic benzotriazoles in blue and green mussels [ng/g lw]. Geometric means in parenthesis.

	UV-32	20	UV-326	UV-326		7	UV-328	
Cambodia	0/2	n.d.	0/2	n.d.	0/2	n.d.	2/2	120 (110)
China	0/5	n.d.	2/5	60 (33)	4/5	84 (65)	3/5	96 (52)
Hong Kong	0/8	n.d.	2/8	91 (18)	6/8	93 (48)	6/8	200 (75)
India	0/3	n.d.	0/3	n.d.	0/3	n.d.	0/3	n.d.
Indonesia	0/2	n.d.	1/2	33 (22)	2/2	58 (45)	2/2	120 (110)
Japan	4/7	33 (13)	7/7	450 (260)	3/7	38 (15)	7/7	120 (93)
Korea	0/17	n.d.	13/17	210 (90)	11/17	100 (56)	16/17	220 (150)
Malaysia	0/4	n.d.	1/4	42 (12)	0/4	n.d.	1/4	24 (14)
Philippines	0/2	n.d.	1/2	120 (50)	2/2	150 (150)	2/2	170 (140)
USA	0/15	n.d.	12/15	130 (79)	11/15	61 (45)	3/15	69 (33)
Vietnam	0/3	n.d.	0/3	n.d.	0/3	n.d.	0/3	n.d.

Analytical results demonstrate ubiquitous contamination and widespread distribution of phenolic benzotriazoles. Levels were comparable to those of PCBs, DDTs and PBDEs. However, spatial variation of the concentrations was often high. Significant correlations were found between the concentrations of several phenolic benzotriazoles, which suggests similar sources and compositions of these compounds in commercial and industrial products. While Kameda et al. (2011) reported correlations of UV-326, -327 and -328 with the polycyclic musk HHCB, such correlations were not always found by Nakata et al. (2012). HHCB is an indicator substance for WWTP effluent. It is concluded that in addition to WWTP effluents there may be point sources or other sources, e.g. road dust, influencing the phenolic benzotriazoles concentrations in mussels.

The authors report that the domestic production and import of UV-327 in Japan decreased dramatically from 2436 tons between 2004 and 2009 to only 3 tons in 2010. They assume that this is due to the availability of an alternative in the Japanese market.

Yanagimoto et al. (Yanagimoto et al., 2011) studied the occurrence of UV-327 and -328 in human adipose tissues collected from Japan (2004-2005, n = 22), South Korea (2005-2006, n = 18), China (2002, n = 12), India (2008, n = 5), Spain (2006, n = 12), Poland (1990, n = 12) and the USA (2003-2004, n = 24). In addition foodstuffs collected from Japan were analyzed for UV-326, -327 and -328 (seafood, meat, eggs, vegetables, dairy products, potatoes, pulses, cereals, fruits, n = 32). Some of the foodstuffs originated from other countries than Japan. GC-HRMS/LRMS was used. All data cited are taken from graphs.

The highest concentrations in human adipose tissue were found In Japan and South Korea. In Japan up to ca. 60 ng/g lw UV-327 were detected in human adipose tissues, in South Korea the concentrations reached ca. 45 ng/g, whereas those in Europe were lower (up to ca. 17 ng/g in Spain, up to ca. 11 ng/g in Poland). Lowest concentrations were observed in the USA (up to ca. 5 ng/g lw). Concentrations of UV-328 were generally lower than those of UV-327: up to ca. 35 ng/g lw in Japan, up to ca. 20 ng/g in South Korea and up to ca. 6 ng/g in Spain, whereas UV-328 was not detected in samples from Poland and only in few samples at low concentrations in the USA (up to ca. 2 ng/g lw). No gender- and age-related differences in concentrations were observed.

In foodstuffs ubiquitous contamination with benzotriazole UV-stabilizers was found. Highest concentrations were detected in seafood (up to ca. 1.2 ng/g ww UV-326, 1.4 ng/g UV-327 and 1.7 ng/g UV-328) and meat (up to ca. 1.5 ng/g ww UV-326, 1.2 ng/g UV-327 and 1.0 ng/g

UV-328). Meat with high concentrations was imported from the USA and Australia. Lower concentrations were detected in vegetables (up to ca. 1.0 ng/g ww UV-326, 0.3 ng/g UV-327 and 0.2 ng/g UV-328) and some fruit (up to ca. 0.5 ng/g ww each UV-326, 327 and 328). In dairy products no benzotriazole UV-stabilizers were found. The estimated daily intake of benzotriazole UV-stabilizers through food consumption was 861 ng/person/d. Contamination was mainly due to meat and vegetables (> 50%), which may imply the transfer of benzotriazole UV-stabilizers from plastic trays and wraps.

By way of a poster Nakata et al. (Nakata et al., 2011) reported temporal trends of UV-327 and -328 in archived marine mammal tissues. In addition temporal trends of UV-326, -327 and - 328 in sediment cores were analyzed. Marine mammals sampled were finless porpoises and striped dolphins from Japanese coastal waters (n = 33). Sediment cores were taken from two sample stations at Tokyo Bay, Japan (n = 12). The sedimentation periods (1930-1999) were determined by 210Pb and the particle fraction < 500 μ m was investigated. All data cited are taken from graphs.

UV-327 and -328 were not detected in blubber samples collected around 1980, but in samples taken in 1990 and later. Maximum concentrations of UV-327 and -328 were ca. 45 ng/g lw and ca. 70 ng/g lw, respectively. An increasing trend is identified for UV-327 as well as UV-328.

Sediment cores showed an increasing temporal trend for UV-326, -327 and -328. Results are presented for two different sampling stations. At both sampling stations concentrations start to rise around 1970. Highest concentrations are found for UV-326 (maximum ca. 17 ng/g dw at station A, ca. 31 ng/g at station B), whereas concentrations of UV-327 and -328 were lower (UV-327 maximum ca. 8 ng/g dw at station A, ca. 4 ng/g at station B, UV-328 ca.10 ng/g at station A, ca. 4 ng/g at station B).

UV-320, -326, -327 and -328 were also detected in road dusts. Samples were collected in December 2010 at 9 stations of Route 57, Kumamoto, with a traffic density of approx. 5,000 to 60.000/d (Nakata Presentation, 2011). All data are taken from graphs.

Concentrations were low for UV-320 (n.d. - ca. 3 ng/g dw), higher for UV-328 (ca.2.5 - ca. 40 ng/g) and UV-326 (ca. 8 - ca. 55 ng/g) and at a single sampling point 116.9 ng/g UV-327 was detected (minimum ca. 8 ng/g dw). Concentrations of UV-320, -326 and -328 correlated with traffic density. The authors conclude that that automobile equipment might be a possible source of benzotriazole stabilizers in the environment.

Based on the data set obtained and the physicochemical properties of benzotriazole UV-stabilizers, the authors conclude that UV-327 will be a candidate of the POP Convention.

Watanabe and Noma (Watanabe and Noma, 2010) performed thermal treatment experiments using pilot-scale equipment and waste containing UV-320 as an input material to determine the destruction behavior of UV-320 and possible formation of UV-327 and NOx.

UV-320 was classified as a "Class I Specified Chemical Substance" under the Chemical Substance Control Law in Japan in 2007, which means that it is comparable in nature and toxicity to POPs (Watanabe and Noma, 2010). Manufacture and import of this substance have to be permitted, only specified uses are allowed and import of certain products specified by cabinet orders is prohibited. Therefore production, import and use of UV-320 have declined in Japan. However, it is still used in some countries, such as Korea and China and in Japan it may still be leached from long-life products. It is expected that incineration may be the predominant method of treatment for wastes containing UV-320.

Concentrations of UV-320 and -327 in "refuse derived fuels" obtained from Japanese municipal solid waste after removing the incombustible materials were 7.1 and 20 μ g/kg, respectively. After treatment in the pilot-scale incinerator with two combustion units, bag filter, activated carbon adsorption tower and wet scrubber concentrations in the flue gas (final exit) were 0.0020 μ g/m³ and 0.0042 μ g/m³ for UV-320 and -327, respectively. Bottom ash contained 0.52 μ g/kg UV-320 and 0.063 μ g/kg UV-327, fly ash 0.36 μ g/kg UV-320 and 0.049 μ g/kg UV-320 and 327 in flue gas, bottom ash and fly ash were of the same order of magnitude as those observed at low input concentrations of UV-320.

UV-320 was destroyed mainly in the primary combustion zone. Overall destruction efficiency of UV-320 in input at a concentration of 5000 mg/kg was > 99.9999%. The input amount of UV-320 did not affect the formation and destruction behavior of UV-327 and NOx.

Other Asian studies:

Kim et al. (Kim et al., 2011b) developed a multiresidue analytical method for the determination of emerging pollutants including UV-234, -320, -326, -327, -328 and -329 in fish. The concentrations in fish muscle tissue were given on a lipid weight (lw) basis and the method detection limits were 0.3 - 9 pg/g for the UV-stabilizers mentioned above. Five individual fish samples belonging to three species of fish from Manila Bay, the Philippines were analyzed. Samples were collected during June 2008. Concentrations ranged from < method detection limit to 179 ng/g lw, suggesting the ubiquitous contamination in Manila Bay.

	bluetail mullet	coral grouper	flathead grey mullet M. cephalus (n=3)				
	V. buchanani (n=1)	E. corallicola (n=1)	mean	Min-Max			
UV-234	not detected	14.3	34.6	22-47.1			
UV-320	9.60	0.78	6.88	4.11-9.15			
UV-326	211	n.d.	18.9	no data given			
UV-327	2.57	18.5	14.6	10.5-18.5			
UV-328	18.4	21.1	105	30.2-179			
UV-329	not detected	39.4	7.29	6.69-7.89			

Table 41: Concentrations of phenolic benzotriazoles in fish muscle tissue [ng/g lw]

Using the same method Kim et al. (Kim et al., 2011c) studied contamination of fish from Manila Bay, the Philippines, with benzotriazole UV-stabilizers including UV-234, -320, -326, - 327, -328 and -329. Manila Bay is one of the pollution hot spots in the seas of East Asia with a very dense population and significant fisheries and aquaculture activities. It serves as a sink and transit area for the domestic and industrial wastes from metro Manila and the surrounding provinces. Many people depend on fish from the bay for food. During January and June 2008 58 fish specimens belonging to 20 species were collected from the local fish markets. Only fishes from Manila Bay were selected and analyzed. The method quantification limits were 1-27 pg/g lw.

Benzotriazole UV-stabilizers were detected, each at ng/g level in almost all fish samples, indicating ubiguitous contamination in coastal waters. Among the 8 targeted substances UV-328 was predominantly found with a mean concentration of 34.2 ng/g lw, implying large scale production and use of this compound in the Philippines. UV-328 was found in 88% of analyzed specimens (n = 58), UV-320 and UV-234 in 79% and 55%, respectively. UV-326, -327 and -329 were detected in less than half of the samples suggesting smaller amount of use or lower bioavailability. Generally concentrations of UV-320, -326, -327 and -328 in fish samples from the Philippines were higher than those reported in marine fish from shallow waters of Japan (Nakata et al., 2009a), which is attributed to large scale usage of the substances and/or the release of untreated wastewater containing the substances. In line with the results of Nakata et al. (2009a) concentrations of UV-320, though frequently detected, were lower than that of UV-234 and -328. According to the authors this may indicate the differences in accumulation and biodegradability of UV-320. Significant positive correlations were found between UV-234 and -328, UV-234 and -329, UV-320 and -327 and UV-320 and UV-328. From this it is suggested that fish in Manila Bay are exposed to benzotriazole UV-stabilizers originating from the same sources which are distributed homogenously in the bay. Examination of the relative contributions of each analyte to the total concentrations of analytes revealed that from the substances relevant for the SVHC dossier UV-328 was predominant. Compositions of the benzotriazole UV-stabilizers were different even in fishes belonging to the same family whereas some composition pattern was observed in fishes belonging to different families. This may be due to different availability, different metabolic capacity or selective uptake of the substances. Concentrations of UV-234, -320, -326, -327, -328 and -329 did not show any relation with fish length and weight. Therefore, differences in accumulation/exposure pattern indicate the species specificity in fish samples. Concentrations measured in the different fish species varied greatly depending on the species within one to two orders of magnitude. This wide variation in concentrations indicates species-specific accumulation and elimination of the substances.

High concentrations of the sum of the investigated 8 substances were found in bumpnose trevally (Caranoides hedlandensis, n = 3), bluetail mullet (adult) (Valamugil buchanani, n = 1), common ponyfish (Leiognathus equulus, n = 3) and coral grouper (adult) (Ephinephelus corallicola, n = 1). These high concentrations (several hundred ng/g lw) indicate that these compounds are preferably accumulated by these species and/or that these species may have low metabolic capacity to eliminate benzotriazole UV-stabilizers. All these fishes belong to the demersal habitat.

Table 42: Concentrations of benzotriazole UV-stabilizers in marine species from Manila Bay, the Philippines

	lipid content [%]	UV-234 [ng/g lw]	UV-320 [ng/g lw]	UV-326 [ng/g lw]	UV-327 [ng/g lw]	UV-328 [ng/g lw]	UV-329 [ng/g lw]	Σ 8 benzotriazole UV-stabilizers
detection frequency [%]		55	79	19	43	88	41	
Min. – Max. in 20 fish species (n = 58)	0.13- 2.61	n.d 126	n.d. – 28.7	n.d 211	n.d 221	n.d 563	n.d 96.7	$6.5 \pm 11.1 - 316 \pm 460$

Kim et al. (Kim et al., 2012) used the same method for determining UV-234, -320, -326, -327 and -328 in house dust from the Philippines. During August 2008 house dust samples were collected from a residential area (Malate, n = 17) and near a large-scale open dumping area of municipal wastes (Payatas, n = 20) in Manila. People live directly at and even on the dumping area (http://www.dr-koelsch.de/html/payatas.html). House dust was collected in separate vacuum-cleaner bags used in each of the sampled house, which consist of dust from living room, kitchen and bedrooms. Dust was not collected from under furniture or in crevices between cushions. Obtained dust samples were combined individually for each house and sieved with a 500 μ m mesh. Data on the details of the house, the possible sources of dust, floor area, number of computers/televisions, furniture and type of flooring were documented in a questionnaire at the time of sample collection.

Table 43: Concentrations of benzotriazole UV-stabilizers in house dust samples from Malate and Payatas in the Philippines

Target	Malate					Payatas				
compounds	DFa [%]	Median [ng/g]	Average [ng/g]	Min. [ng/g]	Max. [ng/g]	DFa [%]	Median [ng/g]	Average [ng/g]	Min. [ng/g]	Max. [ng/g]
UV-234	94	84	148	n.d.b	817	95	41	63	n.d.	212
UV-320	82	4.7	6.6	n.d.	25	65	3.0	6.9	n.d.	75
UV-326	88	50	53	n.d.	275	65	6.2	17	n.d.	133
UV-327	88	19	28	n.d.	73	80	10	10	n.d.	32
UV-328	82	27	50	n.d.	304	85	12	18	n.d.	48
Σ		147	285	n.d.	1020		118	115	n.d.	277

a DF: detection fequency

b n.d. = not detected

UV-234, -320, -326, -327 and -328 were frequently detected indicating ubiquitous contamination of the indoor environments. Among the target compounds, UV-234, -326 and -328 were the predominant compounds. The most abundant was UV-234, with a median value of 84 ng/g in Malate and 41 ng/g in Payatas. Significantly higher concentrations of UV-326 and -327 were found in house dust samples from Malate than those from Payatas, indicating possible differences in usage patterns of household products such as TV, waxes, coating materials, paints etc. between the two locations. Household products are considered the major source of contamination in the indoor microenvironment. The composition of phenolic

benzotriazoles differed among the houses even within the same sampling region. It was not possible to distinguish the sources of the contamination. However, the correlations found for most of the benzotriazole UV-stabiliszers in house dust samples indicate a common source. This is in line with the results from other investigations (Kim et al.2011a, Nakata et al. 2009a)

Generally, levels of benzotriazole UV-stabilizers in dust from the Philippines are comparable to or lower than those measured by Carpinteiro et al. (2010b) in dust from Spain or the USA. Lower levels are attributed to lesser usage of the respective compounds in the Philippines.

Zhang et al. (Zhang et al., 2011) investigated UV-326, UV-327 and UV-328 in surface sediment samples (0-20 cm) collected from rivers in China (6 samples from river Songhua in 2009) and the U.S. (3 samples both from river Saginaw in 2002 and river Detroit in 1998). Five sewage sludge samples were collected from five WWTPs serving large cities located along the Songhua River in China in July 2009. Sediment and sludge samples taken from 4-6 spots within 10 m at a given sampling location were pooled to obtain a representative sample. UV-326, UV-327 and UV-328 were determined by use of a GC-MS.

The limit of detection (LOD) and the limit of quantification (LOQ) for sediment analysed in this study were 0.02 and 0.06 ng/g for UV-327 and 0.1 and 0.33 ng/g for both UV-326 and UV-328. The method LOD and LOQ values for sludge samples were 0.1 and 0.3 ng/g for UV-327 and 0.5 and 1.65 ng/g for both UV-326 and UV-328.

UV-326 was detected in 2 of 6 sediment samples from the Chinese River (1.71 and 2.01 ng/g dw) in 1 of 6 sediment samples from the U.S. (5.88 ng/g dw) and in all 5 sewage sludge samples from China (23.3-136 ng/g dw, mean 77.4 ng/g dw).

UV-327 was detected in 1 of 6 sediment samples from the Chinese River (0.310 ng/g dw) in 3 of 6 sediment samples from the U.S. (0.22-1.90 ng/g dw, mean 0.850 ng/g dw) and in 4 of 5 sewage sludge samples from China (1.80-8.40 ng/g dw, mean 3.68 ng/g dw).

UV-328 was detected in all 6 sediment samples from the Chinese River (2.06 - 7.12 ng/g dw, mean 3.81 ng/g dw) in 5 of 6 sediment samples from the U.S. (0.72-224 ng/g dw, mean 116 ng/g dw) and in all 5 sewage sludge samples from China (40.6-5920 ng/g dw, mean 1300 ng/g dw).

The concentration of UV-328 in sludge was the highest (mean: 1300 ng/g dw) among the target compounds.

Ruan et al. (Ruan et al., 2012) analyzed UV-234, -320, -326, -327, -328, -329 and -350 in municipal sewage sludge in China using an HPLC-MS/MS method. The method quantification limits were from 0.15 (UV-234) to 0.77 (UV-320) ng/g dw. Sixty sewage sludge samples from WWTPs in 33 cities were collected in 2010 and 2011. Most of the WWTPs are located in economically developed provinces in China. Samples were taken from freshly digested sludge at the dewatering process. The most dominant analogue was UV-234 at a median concentration of 116 ng/g dw. The abundance was successively followed by UV-329, -326 and -328 with median concentrations of 66.8, 67.8 and 57.3 ng/g dw respectively. UV-327 and UV-350 had low detection frequency, while UV-320 was not detectable in any sample. According to the authors the observed composition pattern in the sludge samples was quite consistent with the global production volumes of benzotriazole UV-stabilizers (according to the OECD and US EPA HPV databases).

Significant correlations were found among the phenolic benzotriazole concentrations and the daily treatment volume of the WWTPs was moderately correlated UV-329 and UV-328. Results from degradation prediction and multimedia fate simulation based on a quantitative structure-property-relationship (QSPR) model at screening level based on EPISuite and therefore comparable with the simulations done for the presented dossiers implied that the commercial benzotriazole stabilizers and their plausible transformation products might be persistent in the environment.

Table 44:	Concentrations	of	benzotriazole	UV-stabilizers	in	sludge	from	Chinese	municipal
WWTPs									

Analyte	Detection frequency	Concentrations [ng/g dw]	Median [ng/g dw]
UV-234	58/60	0.96 – 235	116
UV-320	0/60	n.d.	-
UV-326	59/60	4.00 - 319	67.8
		two extreme values: 2930 and 3390	
UV-327	24/60	1.53 - 133	14
UV-328	58/60	3.54 - 213	20.6

		one extreme value: 24,700	
UV-329	59/60	0.57 – 757	66.8
UV-350	5/60	1.88 - 42.7	13.8

Australian studies:

Liu et al. (Liu et al., 2011b; Liu et al., 2012) developed a method for simultaneous determination of benzotriazoles and UV-filters (including UV-326 and -329) in ground water and WWTP effluent and biosolid samples using GC-MS/MS. The method was applied to screen the selected substances in samples from Bolivar WWTP in Adelaide, South Australia. The WWTP serves a population of 1,300,000 and is designed to have dry weather flow of 148.5 ML/d. About 75% of the inflow is from domestic sources, 25 % from industrial sources. The WWTP consists of primary sedimentation, secondary activated sludge treatment, stabilization lagoons and dissolved air flotation/filtration. The effluent is piped to a vegetable growing region for irrigation, or recharged into aquifer on site. The sludge line comprises mesophilic anaerobic digestion and sludge stabilization lagoons.

Groundwater samples were collected from an aquifer storage and recovery well at a depth of 300 m below ground within the WWTP site. Biosolid samples were collected from different sludge treated process (sludge is dewatered and dried using a combination of sludge drying lagoons, centrifugation and agitated air drying). 3 parallel samples were collected for each sample type.

In groundwater and effluent water concentrations of UV-326 and -329 were below the limits of quantification (LOQ). The LOQ were: 4.9 ng/L in tap water and 11.0 ng/L in effluent for UV-326 and 18.6 ng/L in tap water and 16.0 ng/L in effluent for UV-329. The concentration in biosolid samples was 49.9 ± 7.4 ng/g for UV-326 (LOQ 1.1 ng/g) and 122.9 ± 7.1 ng/g for UV-329 (LOQ 27.4 ng/g).

Results published in 2012 focus on the removal processes in the WWTP. 24 h composite water samples and samples of sludge (24 h composite or grab) and influent suspended solids were collected in April and October 2010. The average removal efficiencies of suspended solids, BOD5 and NH4-N were above 99% during the sampling periods. The highest value of LOD for the target analytes (4 benzotriazoles and 6 UV-filters including UV-326 and -329), were 16.3 ng/L in the influent, 14.1 ng/L in the effluent and 8.2 ng/g in biosolid samples.

All water and sludge concentrations are taken from graphs. UV-326 was detected in the influent in concentrations of ca. 35 ng/L (April) and ca. 20 ng/L (October), UV-329 in concentrations of ca. 230 ng/L (April) and ca. 420 ng/L (October). According to the authors both substances were completely removed from the water phase. However, removal rates of both > 100% and < 0% were noticed in some treatment stages, which might be due to variations in the input and output concentrations. Concentrations of UV-326 and UV-329 in influent suspended solids were always near 100 ng/g. Both substances are further detected in all other sludge samples taken after different treatment steps.

A mass balance analysis was applied to establish mass flux in the plant and removal mechanisms. However, few data were available, concentrations in water and sludge varied considerably with different treatment stages. The authors discuss plenty uncertainties associated with the mass balance analysis, but nevertheless state that sorption onto sludge played a dominant role in the removal of UV-326 in the WWTP whereas biological degradation played a significant role for UV-329.

American studies investigating the environmental impact of a certain industrial point source:

Jungclaus et al. (Jungclaus et al., 1978) analyzed industrial WWTP effluent and receiving waters and sediments from an American specialty chemicals manufacturing plant producing organic compounds and running a badly performing WWTP. 16 water samples and 19 sediment samples were taken in 1975 and 1976 and the compounds contained were identified, beside others UV-320, -327 and -328. River water and sediments were collected in Providence River and its tributary Pawtuxet River (Pruell et al., 1984).

UV-328 was detected in industrial WWTP effluent (0.55 - 4.7 ppm), in river water (7 - 85 ppb)

and in sediments (1-100 ppm). UV-320 and UV-327 were detected only in sediment, with concentrations of 40 ppm and 2 – 300 ppm, respectively.

Lopez-Avila and Hites (Lopez-Avila and Hites, 1980) investigated transport of pollutants in sediments in the USA. The wastewater from a small specialty chemicals manufacturing plant located on the Pawtuxet River (Rhode Island) contaminated the water and sediment of that river, which flows into the brackish Providence River and Narragansett Bay. UV-328 had been manufactured in the plant since 1970. Wastewater samples from the clarifier tank, water samples and sediment cores were taken. Reported concentrations represent minimum values since they had not been corrected for solvent extraction efficiencies. Average water concentrations for UV-328 (geometric averages of 2-5 values measured at the specified locations at different times) were 3000 ppb in the wastewater of the plant, 40 ppb in river water near the plant, 10 ppb in more distant river water, 8-9 ppb in the mouth of the Pawtuxet River and 0.5-2 ppb in the Providence River. The concentrations follow the rules of simple dilution. UV-327 was manufactured at the plant between 1963 and 1972. It was not detected in any of the water samples.

Eight sediment cores were taken at three locations in the Pawtuxet River. The sites were selected for an abundance of fine-grained material. Further sediment cores were taken at 4 locations in the Pawtuxet Cove and 13 locations in the Providence River and Narragansett Bay. The core concentrations of the compounds in the sediment have been condensed into a single number. However, the authors feel the values given are representative of the sediment concentrations. Concentrations decrease both with depth in the sediment and with increasing distance from the discharge.

Pawtuxet River				Pawtuxet	ixet Providence River			
	near plant	mid river	near dam	Cove	near	far	bay	
UV-327	300	400	20	80	20	2	0.5	
UV-328	300	300	70	100	10	5	0.6	

Table 45: Concentrations of phenolic benzotriazoles in sediment cores (ppm)

Pruell et al. (Pruell et al., 1984) developed an analytical method for the determination of PAHs and phenolic benzotriazoles in clams. Concentrations of UV-327 and -328 were measured in hard shell clams (Mercenaria mercenaria) purchased from Rhode Island seafood stores in 1979. Personnel in nine of the 13 stores surveyed indicated that the clams were harvested from Narragansett Bay. Three seafood stores were sampled a second time to determine if the higher values obtained at these establishments were representative of their usual stock. As controls, clams were collected from a relatively unpolluted site in lower Narragansett Bay. The detection limit for specific compounds was ca. 0.1 ng/g ww.

The levels in purchased clams were generally higher than the concentrations found in clams collected from a lower Narragansett Bay control location. However, also in control samples both substances were detected. In summary UV-328 and UV-327 were present in clam tissue in concentrations ranging from 7 – 65 ng/g ww and from 1.0 - 8.5 ng/g ww (including controls). The ratio of UV-328 to UV-327 in clams varied from 2.7 to 9.5. This is similar to the ratio in surface sediments of the bay which ranges from 2.0 to 7.6. A significant correlation existed between UV-327 and UV-328.

Reddy et al. (Reddy et al., 2000) examined the free and bound fractions of different substituted benzotriazoles in sediment cores from the Pawtuxet River and Narragansett Bay in the U.S. The chosen benzotriazoles were produced from 1961 to 1985 by a major chemical plant located on the Pawtuxet River. Beside others, UV-326, -327 and -328 were investigated. Previous research has used these compounds as specific tracers of inputs from the Pawtuxet River into Narragansett Bay sediments and they are highly enriched in the sediments of both.

The Pawtuxet River sediment core was collected in 1989 and sectioned at 2-3 cm intervals. Eleven sections from 0-2 cm to 50-52 cm were analyzed. The sedimentation rates in this section of the river are 2-3 cm/year. The redox discontinuity, determined visually, was in the top 2 cm of the core. The Narragansett Bay core was collected in 1997. Six sections from the top 13 cm of the core were analyzed. The sediments in this area become anoxic within a few millimeters of the surface and have a sedimentation rate of about 0.3 cm/year. The deepest sections of both cores were the approximate depths of where the phenolic benzotriazoles were

no longer detected and should roughly be equivalent to the initial date of production of these compounds (1961-1979). The method detection limit was ca. 20 ng/g for each (free and bound) fraction.

In the Narragansett Bay core UV-327 and -328 were detected at trace levels in the 10-13 cm section and their concentrations generally increased up-core (with concentrations as high as 25 μ g/g). UV-326 was detected at much lower concentrations. UV-327 and -328 were not detected in the bound fraction in the Narragansett Bay core.

In the Pawtuxet River core all benzotriazoles were detected in the free fraction. UV-327 was most abundant: the highest concentration was ca. 5 mg/g and it was observed down to 50-52 cm. The other benzotriazoles were only present in the top 20 cm of the core. UV-326 and -327 were also found in the bound fraction of the Pawtuxet River core in at least the top 15 cm. However, the maximum percentage bound was 0.04%.

Benzotriazoles that had alkyl substitution in ortho position to the hydroxyl group were less likely to be found in the operationally defined bound fraction than compounds that did not have this substitution.

Hartmann et al. (2005) took sediment cores at three locations in Narragansett Bay in 1997 (Apponaug Cove, Seekonk River, Quonset Point). The cores were analyzed for several contaminants including UV-327 and UV-328. The phenolic benzotriazoles were used as markers indicating the years of their introduction (1963 for UV-327 and 1970 for UV-328). Two of the cores were split into 2 cm sections, and the third core (Quonset Point) was split into 10 cm sections.

Sharp breaks in the concentrations of UV-327 and UV-328 marking their introduction were successfully used to determine the sedimentation rate at Quonset Point. Both the Quonset Point and Seekonk River cores had subsurface maximums for phenolic benzotriazoles, which were consistent with expected inputs to the environment. The Apponaug Cove core showed an increase of the contaminants at the surface indicating a recent event in which more contaminated sediments were deposited at that location. The distributions of phenolic benzotriazoles at Apponaug Cove and in the Seekonk River indicate that there was a disturbance in the depositional environment relative to cores collected at these locations in 1986, demonstrating the potential for buried contaminants to be remobilized in the environment even after a period of burial.

At Quonset Point the phenolic benzotriazole profile increased down core through the 40-50 cm section before decreasing in the 50-60 cm section. Below the 50-60 cm section, UV-327 and UV-328 were below the detection limit of 10 ng/g dw. In the 50-60 cm section UV-327 is much more prominent than UV-328. Moving up core, UV-328 progressively accounts for more of the sum of both phenolic benzotriazoles. This reflects the earlier introduction (1963) and subsequent earlier discontinuation (1972) of UV-327 relative to UV-328 (1970 and 1985, respectively).

At Apponaug Cove surface concentrations were higher than the lower sections of the core. There could be degradation in the oxic surface layer of the sediments with subsequently lower concentrations in the deeper sections. However, data from a core taken in 1986 had a profile more consistent with the appearance of the different analytes. Therefore the authors assume that the distribution of phenolic benzotriazoles represents resuspended sediment transport and deposition of materials with high concentrations.

Data from the Seekonk River core also show high concentrations in the surface layer. Another core taken in the same area in 1986 showed a more orderly decrease down to 70 - 80 cm. The authors assume that some sedimentary layers were removed. Additional evidence of a disturbance is found in the ratio of the phenolic benzotriazoles. The lowest core section with phenolic benzotriazoles (12 - 14 cm) should have high ration of UV-327 to UV-328 due to their production history, but in this case actually had a lower ratio of UV-327 to UV 328 than the sections above it.

Quonset Poi	int core		Apponaug C	ove core		Seekonk Riv	ver core
depth	UV-327	UV-328	depth	UV-327	UV-328	UV-327	UV-328
[cm]	[ng/g dw]	[ng/g dw]	[cm]	[ng/g dw]	[ng/g dw]	[ng/g dw]	[ng/g dw]
0 - 2	ca. 40	ca. 160	0 - 2	ca. 130	ca. 270	ca. 30	ca. 120
0 - 10	ca. 60	ca. 260	2 - 4	ca. 30	ca. 80	ca. 20	ca. 70
10 - 20	ca. 80	ca. 360	6 - 8	ca. 50	ca. 140	ca. 30	ca. 140
20 - 30	ca. 100	ca. 840	10 - 12	ca. 70	ca. 120	-	-
30 - 40	ca. 130	ca. 1100	12 - 14	-	-	ca. 5	ca. 20
40 - 50	ca. 690	ca. 1180	20 - 22	n.d.	n.s.	n.d.	n.d.
50 - 60	ca. 480	ca. 40	30 - 32	n.d.	n.d.	-	-
60 - 70	n.d.	n.d.	38 - 40	-	-	n.d.	n.d.
80 - 90	n.d.	n.d.	40 - 42	n.d.	n.d.	-	-
100 - 110	n.d.	n.d.	48 - 50	-	-	n.d.	n.d.
119 - 129	n.d.	n.d.					
nd = not de	toctod						

Table 46: Concentrations of phenolic benzotriazoles in sediment cores from Narragansett Bay (concentrations taken from a graph)

n.d. = not detected

not measured

At Apponaug Cove the phenolic benzotriazole profile indicates a much higher surface concentration than the lower sections of the core. Because the production of UV-328 was discontinued 12 years before the core was taken and the production of UV-327 25 years before that date, the authors attribute the high surface concentrations to resuspended sediment transport and deposition of materials in Apponaug Cove with relatively high concentrations of phenolic benzotriazoles. The ratio of UV-327 to UV-328 also increases in the surface section and may indicate a disturbance of older sediments having higher UV-327 levels.

ANNEX 7: Available Information on Endocrine Disrupting properties of phenolic benzotriazoles

In-vitro-Studies

The estrogenic activity of several phenolic benzotriazoles was tested in a Yeast-Estrogen-Screen-assay (YES-assay) with human estrogenic receptors. In the study of Miller et al. (Miller et al., 2001) UV-327 and UV-329 (CAS 3147-75-9) were tested and in the study of Kawamura et al. (Kawamura et al., 2003) UV-327, UV-234 (CAS 70321-86-7), UV-326 (CAS 3896-11-5), UV-328 and UV-P (CAS 2440-22-4). Both studies showed that none of the phenolic benzotriazoles tested was triggering an estrogenic receptor activity.

In a study of Kunz et al. (Kunz et al., 2006) UV-360 (CAS 103597-45-1) was tested in a Yeast-Estrogen/Androgen-Screening-assay (YES/YAS-assay). No effects were reported.

In-vivo-Studies

In a recent review of the U.S. National Toxicology Program on the phenolic benzotriazoles UV-P, UV-329, UV-326, UV-320, UV-327, UV-328, UV-234, UV-360 as well as CAS 84268-36-0 (i.e. M1), 84268-33-7 (i.e. the methyl ester of M1), 84268-08-6 (i.e. a more complex ester of M1) and CAS 104810-48-2/104810-47-1 (i.e. an oligomeric ester of M1) (National Institute of Environmental Health Sciences, 2011) an overview over the available toxicity studies on mammals is given. There are several indications on effects mentioned that might be caused by endocrine disruption, e.g. reduced concentrations of testosterone, higher concentrations of CYP450, or higher activity of ethoxyresorufin-O-deethylase (EROD-activity). As in these cases there are also indications for toxic effects on the liver reported, the effects might actually be only secondary effects. With the present knowledge it is not possible to attribute them unambiguously as endocrine adverse effects.

Preliminary assessment of ED-properties for the phenolic benzotriazoles

There are several indications on effects of phenolic benzotriazoles mentioned that might be caused by endocrine disruption, e.g. reduced concentrations of testosterone, higher concentrations of CYP450, or higher activity of ethoxyresorufin-O-deethylase (EROD-activity). As in these cases there are also indications for toxic effects on the liver reported, the effects might actually be only secondary effects. With the present knowledge it is not possible to attribute them unambiguously as endocrine adverse effects of an equivalent level of concern.

ANNEX 8: Abbreviations

°C	Degrees centigrade
Å	Angstrom
avg.	Average
В	Bioaccumulative
BAF	Bioaccumulation factor
BCF	Bioconcentration factor
BMF	Biomagnification factor
CAS	Chemical Abstracts Service
CLP	Classification, labelling and packaging (of substances and mixtures)
C&L	Classification and labelling
cm	Centimetres
cm³ CMR	Cubed centimetres
CYP450	Carcinogenic, mutagenic, toxic to reproduction Cytrochrome P 450
d	Day
DDT	Dichlorodiphenyltrichloroethane
DegT50	Time interval after which 50% of a substance is degraded
DF	Detection frequency
DT50	Time interval after which 50% of a substance is degraded or
	disappeared otherwise from the test medium
DisT50	Time interval after which 50% of a substance disappeared from the
	test medium (no degradation)
dw	Dry weight
EC	European Community
ECHA EPA	European Chemicals Agency
EROD	Environmental Protection Agency Ethoxyresorufin-O-deethylase
EU	European Union
g	grammes
GC-MS/MS	Gas chromatography – tandem mass spectrometry
GC-HRMS/LRMS	Gas chromatography – high resolution mass spectrometry/low
-	resolution mass spectrometry
GLP	Good laboratory practice
h	Hour
H351	Classification: suspected of causing cancer
H373	Classification: May cause damage to organs through prolonged or
11412	repeated exposure
H412 HHCB	Classification: Harmful to aquatic life with long lasting effects 1,3,4,6,7,8-Hexahydro-4,6,6,7,8,8-hexamethyl-cyclopenta-[g]-2-benzo-
ппсв	pyrane, a polycyclic musk, CAS 1222-05-5
HPLC	High performance liquid chromatography
HPLC-MS/MS	High performance liquid chromatography – tandem mass
	spectrometry
IUPAC	International Union of Pure and Applied Chemistry
k	Rate constant (e.g. for biodegradation in sewage treatment plants)
Kair-water	Air-water partition coefficient
Kg	Kilograms
Кос	Organic carbon-water partition coefficient
Kow	Octanol/water partition coefficient (log value)
Kp KPa	Partition coefficient Kilopascals
L (or l)	Litres
LC	Liquid chromatography
LC-MS	Liquid chromatography – mass spectrometry
LC-MS/MS	Liquid chromatography – tandem mass spectrometry
·	

LC50	Lethal concentration for 50% of the test organisms	
LOD	Limit of detection	
LOQ	Limit of quantification	
lw	Lipid weight	
M	Molar	
m ²	Metres squared (area)	
m ³	Cubed metres (volume)	
Max	Maximum	
Min	Minimum	
MITI	Ministry of International Trade and Industry (Japan)	
mg	Milligrams	
ml ML	Millilitres Megalitre	
Mol	Moles	
Mmol	Millimoles	
MS	Mass spectrometry	
μg	Micrograms	
n	Number (e.g. number of samples)	
n.d.	Not detected	
NER	Non-extractable residues	
NITE	National Institute of Technology and Evaluation, Japan	
nm	Nanometres	
NOEC	No-observed effect concentration	
ос	Organic carbon	
OECD	Organisation for Economic Co-operation and Development	
Р	Persistent	
Pa	Pascals	
PBDE	Polybromodiphenyl ether	
PBT	Persistent, bioaccumulative and toxic	
PCB	Polychlorinated biphenyl	
POP	Persistent organic pollutant	
PPB PPM	Parts per billion Parts per million	
QSAR	Quantitative structure-activity relationship	
QPREF	QSAR Prediction Reference Format	
QSPR	Quantitative structure-property-relationship	
REACH	Registration, Evaluation, Authorisation and restriction of	
	Chemicals Regulation (EC 1907/2006)	
S	Seconds (time)	
SIM	Selective ion monitoring	
STOT-RE	Specific target organ toxicity – repeated exposure	
SVHC	Substances of very high concern	
Σ	Sum	
Т	Toxic (hazard classification)	
US or USA	United States of America	
UV	Ultraviolet	
UV-234	A phenolic benzotriazole UV stabilizer, CAS 70321-86-7	
UV-320	2-benzotriazol-2-yl-4,6-di-tert-butylphenol, CAS 3846-71-7	
UV-326	A phenolic benzotriazole UV stabilizer, CAS 3896-11-5	00 1
UV-327 UV-328	2,4-di-tert-butyl-6-(5-chlorobenzotriazol-2-yl)phenol,CAS 3864- 2-(2H-benzotriazol-2-yl)-4,6-ditertpentylphenol,CAS 25973-55-1	
UV-328 UV-329	A phenolic benzotriazole UV stabilizer, CAS 3147-75-9	L
UV-350	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec-butyl)phenol,CAS	
0.6-220	36437-37-3	
UV-360	A phenolic benzotriazole UV stabilizer, CAS 103597-45-1	
UV-571	A phenolic benzotriazole UV stabilizer, CAS 105557 45 1	
UV-928	A phenolic benzotriazole UV stabilizer, CAS 73936-91-1	
UV-P	A phenolic benzotriazole UV stabilizer, CAS 2440-22-4	
vB	Very bioaccumulative	
		107

Very persistent
Very persistent, very bioaccumulative
When applicable
Wet weight
Waste water treatment plant
Yeast-estrogen-screen
Yeast-Estrogen/Androgen-Screening